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### (57) Abstract

A promoter comprising the DNA sequence of an oil seed rape cysteine protease gene promoter of class 1, 2 or 6 is described. The promoter may be used in an expression system for at least the tissue or tissues of a germinating seedling or developing grain or plant (e.g. in the root, cotyledons, leaves and stem). In a preferred embodiment, the expression system comprises a disrupter gene fused to a promoter according to the present invention.

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# CYSTEINE PROTEASE PROMOTER FROM OIL SEED RAPE AND A METHOD FOR THE CONTAINMENT OF PLANT GERMPLASM

The present invention relates to promoters and to a construct comprising the same.

The present invention also relates to a method for the containment of plant germplasm.

In particular, the present invention relates to the use of a promoter for the expression of a gene of interest (GOI) in a specific tissue or tissues of a plant.

More particularly, the present invention relates to promoters for cysteine proteases. The present invention also relates to the application of these cysteine protease promoters to express a GOI in a specific tissue or tissues of a plant.

Promoters control the spatial and temporal expression of genes by modulating their level of transcription. Early approaches to genetically engineered crop plants utilised strong constitutive promoters to drive the expression of foreign genes. As strategies in plant biotechnology have become more sophisticated, there are requirements for specific promoters to target transgene expression to a particular tissue or to a particular developmental stage.

Cysteine proteases are members of a large multigene family in plants (Praekelt et al., 1988; Goetting-Minesky and Mullin, 1994), animals (Wiederanders et al., 1992) and protozoa (Mallinson et al, 1994). Cysteine proteases are synthesised as an inactive precursor (Praekelt et al., 1988). The pre-pro-enzyme is targeted to the secretory pathway (Marttila et al., 1995) and post-transcriptionally processed in the vacuoles by proteolytic cleavage of the propeptide fragment to produce the active enzyme (Hara-Nishimura et al., 1993 and 1994).

Plant cysteine proteases participate in different metabolic events of physiological importance. During seed germination and plant senescence they are involved in protein degradation (Jones et al., 1995; Valpuesta et al., 1995; Smart et al., 1995) and play a key role in protein storage mobilisation during germination (Boylan and Sussex, 1987). During seed development, cysteine proteases catalyse the post-translational processing of protein precursors into their mature form (Hara-Nishimura et al, 1995). In addition, some are subjected to hormonal regulation either by giberellic acid (Koehler and Ho, 1990; Watanabe et al., 1991) or ethylene (Cervantes et al., 1994; Jones et al., 1995). Others are induced in response to stress like wounding (Linthorst et al., 1993; Lidgett et al., 1995), dehydration (Guerrero et al., 1990), cold (Schaffer and Fischer, 1988) or are implicated in plant-microbe interactions (Goetting-Minesky and Mullin, 1994).

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Germination specific cysteine proteases have been characterised for barley (Marttila et al., 1995), rice (Watanabe et al., 1991), maize (Debarros and Larkins, 1994), chick-pea (Cervantes et al., 1994), vetch (Becker et al, 1994) and a cysteine protease has been described for oil seed rape (Comai and Harada, 1989). However, the published data for oil seed rape is contradictory. Furthermore, this species is difficult to study due to its amphidiploid nature. Rather than using more conventional and laborious techniques like subtractive or differential screening of cDNA libraries or differential display techniques, potentially generating clones of unknown identity, cysteine proteinases (cysteine proteases) in oil seed rape were studied which are expressed in germinating seeds. Promoters from genes which are uniquely expressed following seed germination were isolated and characterised.

Thus, according to a first aspect of the present invention, there is provided an oil seed rape cysteine protease gene promoter of class 1, 2 or 6.

According to a second aspect of the present invention, there is provided a promoter comprising at least part of a sequence as shown in Figures 19, 20 or 21, or at least part of a sequence that has substantial homology therewith, or a variant thereof.

According to a third aspect of the present invention, there is provided a promoter having the characteristic motifs or features of promoters of the present invention.

According to a fourth aspect of the present invention, there is provided a recombinant DNA construct comprising the promoter as defined above operably linked to a gene which codes for a protein of interest.

According to a fifth aspect of the present invention, there is provided a recombinant DNA construct functional in a plant comprising a disrupter gene encoding a product capable of disrupting cell function, and a promoter as defined above, the disrupter gene being functionally linked to and controlled by an externally regulatable gene control region which includes a promoter which is inducible by the external application of a chemical inducer.

According to a sixth aspect of the present invention, there is provided DNA comprising at least part of the sequence shown in Figures 12, 13, 14, 15, 16 or 17, or at least part of a sequence that has substantial homology therewith or a variant thereof, and which codes for a cysteine protease.

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According to a seventh aspect of the present invention, there is provided a recombinant DNA construct functional in a plant comprising the DNA as defined above operably linked to a promoter.

According to an eighth aspect of the present invention, there is provided an expression system for the tissue or tissues of a plant material, the expression system comprising a gene of interest fused to a gene promoter as defined above wherein the expression system is capable of being expressed in the tissue or tissues of the plant material.

According to a ninth aspect of the present invention, there is provided an expression system comprising a construct as defined above.

According to a tenth aspect of the present invention, there is provided a recombinant plant genome comprising a promoter as defined above, DNA as defined above, a recombinant DNA construct as defined above or an expression system as defined above.

According to an eleventh aspect of the present invention, there is provided a plant, plant seed or plant cell having a recombinant plant genome as defined above.

According to a twelfth aspect of the present invention, there is provided protected germplasm comprising a recombinant DNA construct as defined above.

According to a thirteenth aspect of the present invention, there is provided a plant or seed which is capable of growing to maturity comprising a recombinant DNA construct as defined above.

According to a fourteenth aspect of the present invention, there is provided the use of a gene promoter as defined above to induce expression of a gene of interest when fused to the gene promoter in the tissue or tissues of a plant material.

Preferably, the inducible promoter of the recombinant DNA construct is functionally linked to and controls a repressor protein and the disrupter gene promoter includes an operator sequence which is recognised by the repressor protein, so that in the presence of the inducer the repressor protein is produced which interacts with the operator sequence thereby disabling the second promoter and inhibiting expression of the disrupter gene.

Preferably, the disrupter gene is a nucleotide sequence, which is in sense orientation to an endogenous plant gene which is essential to plant development or a gene conferring a

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desired characteristic on the plant, or comprises a partial sense sequence of the endogenous plant gene.

Preferably, the disrupter gene is a nucleotide sequence which is in antisense orientation to an endogenous plant gene which is essential to plant development or a gene conferring a desired characteristic on the plant.

Preferably, the endogenous plant gene is essential to seed germination or early seedling development.

Preferably, the externally regulatable gene control region is a chemically inducible gene promoter sequence from the glutathione S-transferase system (which is the subject of our International Patent Application No. PCT/GB96/02116), the *Alc* system (which is the subject of our International Patent Application Nos. PCT/GB96/01883 and PCT/GB96/01846) or the ecdysone system (which is the subject of our International Patent Application No. PCT/GB96/01195).

Preferably, the repressor protein gene encodes a bacterial repressor such as the *lac* repressor or a repressor used by 434, P22 or lambda-bacteriophages.

Preferably, the disrupter gene or disrupter promoter contains a "pseudo-operator".

Preferably, the disrupter gene is a cytotoxic gene.

Preferably, the disrupter gene encodes a recombinase or a transposase adapted to excise a nucleotide sequence flanked by recombinase recognition sequences.

Preferably, the recombinant DNA construct is capable of being expressed in the tissue or tissues of a germinating seedling or a developing grain or a plant when the construct is integrated, preferably stably integrated, within the grain's or seedling's or plant's genomic DNA.

Preferably, the expression system is for at least the tissue of a germinating seedling or developing grain or plant (eg in the root, cotyledons, leaves and stem).

Preferably, the expression system is integrated, preferably stably integrated, within a germinating seedling's genomic DNA or a developing grain's genomic DNA or a plant's genomic DNA.

Preferably, the gene promoter is used to induce expression of a gene of interest when fused to the gene promoter in at least the tissue or tissues of a germinating seedling or a developing grain or a plant (eg in the root, cotyledons, leaves and stem).

According to a preferred embodiment of the present invention, the promoter comprises a DNA sequence corresponding to that of the promoter region of the clone pKS12p6, as shown in Figure 19.

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According to another preferred embodiment of the present invention, the promoter comprises a DNA sequence corresponding to that of the promoter region of the clone pKS25p7, as shown in Figure 20.

According to a further preferred embodiment of the present invention, the promoter comprises a DNA sequence corresponding to that of the promoter region of the clone pKS66p1, as shown in Figure 21.

An even more preferred embodiment of the present invention is a seedling, grain or plant comprising a construct comprising a disrupter gene fused to a cysteine protease promoter, wherein the construct is integrated, preferably stably integrated within the seedling's, grain's or plant's genomic DNA, wherein the promoter comprises at least part of a sequence shown in Figures 19, 20 or 21, or at least part of a sequence that has substantial homology therewith, or a variant thereof and wherein the disrupter gene is a gene which encodes barnase ribonuclease.

An even more preferred embodiment of the present invention is a seedling, grain or plant comprising a construct comprising a disrupter gene fused to a cysteine protease promoter, wherein the construct is integrated, preferably stably integrated within the seedling's, grain's or plant's genomic DNA, wherein the promoter comprises at least part of a sequence shown in Figures 19, 20 or 21, or at least part of a sequence that has substantial homology therewith, or a variant thereof and wherein the disrupter gene is a gene which encodes a recombinase adapted to excise a nucleotide sequence flanked by recombinase recognition sequences.

Thus, according to a highly preferred embodiment of the present invention, there is provided a recombinant DNA construct for insertion into the genome of a plant to impart control of plant development thereto, comprises, in sequence:

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- (a) an inducible gene promoter sequence responsive to the presence or absence of an exogenous chemical inducer;
- (b) either a gene encoding a repressor protein under control of the said inducible gene promoter sequence or a gene encoding an inhibitor of the product of the disrupter gene specified at (e) below;
  - (c) an operator sequence responsive to the said repressor protein;
  - (d) a gene promoter sequence of the present invention; and,
  - (e) a gene encoding a protein disrupter of a plant characteristic essential to the growth of the plant, whereby the presence or absence of the exogenous chemical inducer enables either growth to maturity or causes growth to slow down or stop at an appropriate stage.

The inducible promoter used in the present invention may promote expression of the repressor protein in response to stimulation by an exogenous chemical inducer whereby in the absence of the chemical inducer no repressor protein is expressed to interact with the operator thus permitting expression of the disrupter protein gene and in the presence of the chemical inducer repressor protein is expressed thereby preventing expression of the gene encoding the inhibitor of plant development permitting unimpeded plant growth.

The term "plant material" includes a developing caryopsis, a germinating caryopsis or grain, or a seedling, a plantlet or a plant, or tissues or cells thereof, such as the cells of a developing caryopsis or the tissues of a germinating seedling or developing grain or plant (eg in the root, leaves and stem).

The term "gene of interest" or "GOI" with reference to the present invention means any gene of interest. A GOI can be any gene that is either foreign or natural to the plant in question, except for the wild type functional gene when in its natural environment.

Typical examples of a GOI includes genes encoding for proteins and enzymes that disrupt cell function. For example, the gene may be a cytotoxic gene. Alternatively the gene may encode a recombinase, transposase, or a related enzyme with similar properties, adapted to inhibit an endogenous plant gene which is essential to plant development or a gene conferring a desired characteristic on the plant.

A recombinase is an enzyme that recognises a specific excision sequence or set of specific excision sequences and effects the removal of, or otherwise alters, DNA between

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specific excision sequences. Recombinase systems such as the Cre-lox, the FLP, SR1 and SSV1-encoded integrase systems may be used in the present invention.

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Other examples of a GOI include defensive or protective genes, such as genes giving herbicide, fungal or insect resistance. Such genes may be expressed during germination of seedlings at which time the seedlings are particularly vulnerable. Preferably, the gene encodes a protein which confers resistance to biotic and environmental stresses on a plant.

Also included are endogenous genes such as genes encoding  $\beta$ -tubulin and adenine nucleotide translocator (ANT).

The term "disrupter gene" is a gene which, when expressed or repressed specifically at a suitable stage of plant development, will lead to the failure of a plant to reach maturity and to set seed. The origin of the disrupter genes can be from a variety of naturally occurring sources eg human cells, bacterial cells, yeast cells, plant cells, fungal cells, or they can be totally synthetic genes which may be composed of DNA sequences, some of which may be found in nature, some of which are not normally found in nature or a mixture of both. The disrupter genes will preferably be targeted to an essential biochemical function, such as DNA and RNA metabolism, protein synthesis, and other metabolic pathways.

In a preferred embodiment, the disrupter gene is a gene which encodes barnase ribonuclease,  $\beta$ -tubulin or adenine nucleotide translocator (ANT).

The term "variant thereof" with reference to the present invention means any substitution of, variation of, modification of, replacement of, deletion of or the addition of one or more nucleic acid(s) from or to the promoter sequence providing the resultant sequence is capable of expressing a GOI. The term also includes sequences that can substantially hybridise to the promoter sequence. The term also includes DNA which hybridises to the DNA of the present invention and which codes for at least part of a cysteine protease promoter. Preferably, such hybridisation occurs at, or between, low and high stringency conditions. In general terms, low stringency conditions can be defined as 3 x SSC at about ambient temperature to about 65°C, and high stringency conditions as 0.1 x SSC at about 65°C. SSC is the name of a buffer of 0.15M NaCl, 0.015M trisodium citrate. 3 x SSC is three time as strong as SSC and so on.

The term "substantial homology" covers homology with respect to at least the essential nucleic acids/nucleic acid residues of the promoter sequence providing the

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homologous sequence acts as a promoter eg a cysteine protease promoter which is capable of expressing a GOI. Typically, homology is shown when 60% or more of the nucleotides are common with the promoter sequence of the present invention, more typically 65%, preferably 70%, more preferably 75%, even more preferably 80% or 85% and, especially preferred, are 90%, 95%, 98% or 99% or more homology.

The term "construct" - which is synonymous with terms such as "cassette". "hybrid" and "conjugate" - includes a GOI directly or indirectly attached to the promoter of the present invention, such as to form a cassette. An example of an indirect attachment is the provision of a suitable spacer group such as an intron sequence, intermediate the promoter and the GOI. The same is true for the term "fused" in relation to the present invention which includes direct or indirect attachment.

The term "expression system" means that the system defined above can be expressed in an appropriate organism, tissue, cell or medium. In this regard, the expression system of the present invention may comprise additional components that ensure to increase the expression of the GOI by use of the gene promoter.

The DNA of the present invention may be genomic DNA which is in an isolated form and is, preferably, operably linked to a sequence with which it is not naturally associated, or the DNA may be synthetic DNA or cDNA.

Young seedlings at germination represent a vulnerable stage of plant development. Strategies to improve crop production may include expression of genes during this stage to enhance the seedling's resistance to biotic and environmental stresses such as resistance to cold, salt, heavy metals, fungal attack. Protecting seedlings by expressing proteins which provide at least some measure of resistance/tolerance against such stresses, for example antifungal proteins (Cammue et al., 1992; Terras et al., 1993) under the control of a germination specific promoter will limit the expression to a precise phase of development, when the proteins will be most effective. Such development specific expression has further advantages such as avoiding expression of these genes in plant material entering the food chain.

The promoters of the present invention may also be advantageously used in plant germplasm containment systems.

Agriculture uses many crop plants for the production of food for human consumption, for commercial processes yielding products for human consumption, for animal feedstuff

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production, for the development of industrial products and other purposes. The process involves the planting by the farmer of seed which usually has been purchased from a seed producer. The product produced by the crop, be it the whole plant, the seed or fruit of the plant, is harvested and is then used for the various food applications mentioned above.

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The supplied hybrid or inbred seed may incorporate novel genetic information introduced by transformation of the crop giving novel agronomic features such as tolerance to herbicides, insect pests, and fungal diseases, improved yield and/or quality of the harvested product, and novel mechanisms for the control of plant fertility. Such improvements which are made possible through biotechnological research, improve the quality of the plant breeding and improve the agronomic performance of the seed supplied to the farmer.

A problem addressed by the present invention is the containment of crop plants within the area of cultivation. Seeds of cultivated crop plants may be conveyed outside the defined growing area by a number of routes (by birds or small mammals or simply by being dropped during post-harvest transport of a seed crop) where they assume the status of weeds, or they may remain as volunteers in a subsequent crop in later years. It would clearly be appropriate, if it were possible, that cultivated crops be confined to the growing area and prevented from persisting in the wild. It will be appreciated that the problems of crop non-confinement mentioned above become more acute where transgenic crops are involved.

In the same way, pollen can travel long distances through wind and/or insect transportation (dispersion) and remain viable for a long period. Since interspecific crossing between crop plants and their non cultivated related species is possible, a second concern is the escape of pollen from transgenic crops e.g. herbicide resistant crops, to their related weeds species (Mikkelsen et al., 1996). Ways to reduce viability of such hybrids would limit the risk of transgene escape to non-crop species thus avoiding the spreading of plants with enhanced invasiveness or weediness.

It will be appreciated that the use of the seedling promoters of the present invention restricts expression of the disrupter protein gene to a suitable stage of plant development, and also means that it is not necessary to continue to apply an inducer chemical to the plant throughout its lifetime in order to maintain its viability. This has both economic and ecological benefits.

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The invention also provides a genetically transformed plant and parts thereof, such as cells protoplasts and seeds, having incorporated, preferably stably incorporated, into the genome the construct of the present invention.

Thus, the invention provides a plant which can be reversibly inhibited at an appropriate developmental stage in which said plant contains, preferably stably incorporated in its genome, the recombinant DNA construct defined above.

Expression of a protein encoded by a gene is controlled by the interaction of certain regulatory proteins, known as DNA-binding proteins, with a region located upstream of the gene. Within the promoter region, there are located several operator regions which contain a specific oligonucleotide sequence to which these DNA-binding proteins specifically bind. These proteins can lead either to activation or repression of gene expression. Thus, they control the regulated expression of genes.

These DNA-binding proteins, which may in fact be either repressors or activators of gene expression, are herein referred to for the sake of simplicity as "repressors".

The present invention makes use of the well-characterised interaction between bacterial operators with their repressors to control the expression of the disrupter gene function. Bacterial repressors, particularly the *lac* repressor, or repressors used by 434, P22 and lambda bacteriophages can be used to control the expression in plant cells very effectively.

A second operator/repressor system is the subject of our published International Patent Application No. WO90/08827 which is incorporated herein by reference.

A third approach for the down-regulation of the disrupter genes which can be considered is the use of either "antisense", "sense" or "partial sense" technology.

A DNA construct according to the invention may be an "antisense" construct generating "antisense" RNA or a "partial-sense" construct (encoding at least part of the functional gene product) generating "sense" RNA.

"Antisense RNA" is an RNA sequence which is complementary to a sequence of bases in the corresponding mRNA: complementary in the sense that each base (or the majority of bases) in the antisense sequence (read in the 3' to 5' sense) is capable of pairing with the corresponding base (G with C, A with U) in the mRNA sequence read in the 5' to 3' sense.

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Such antisense RNA may be produced in the cell by transformation with an appropriate DNA construct arranged to generate a transcript with at least part of its sequence complementary to at least part of the coding strand of the relevant gene (or of a DNA sequence showing substantial homology therewith).

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"Sense RNA" is an RNA sequence which is substantially homologous to at least part of the corresponding mRNA sequence. Such sense RNA, or partial sense RNA, may be produced in the cell by transformation with an appropriate DNA construct arranged in the normal orientation so as to generate a transcript with a sequence identical to at least part of the coding strand of the relevant gene (or of a DNA sequence showing substantial homology therewith). Suitable sense, or partial sense constructs may be used to inhibit gene expression (as described in International Patent Publication WO91/08299).

Antisense RNA constructs which may be used to down-regulate disrupter genes include those encoding adenine nucleotide translocator.

Other approaches which are, or become, available may also be used.

Further details on such crop containment systems can be found in our published International Patent Application No. WO94/03619 which is incorporated herein by reference.

The promoter of the present invention, then, when linked to an exogenous or foreign gene and introduced into a plant by transformation, provides a means for the regulation of expression of that foreign gene. The method employed for transformation of the plant cells is not especially germane to this invention and any method suitable for the target plant may be employed. Transgenic plants are obtained by regeneration from the transformed cells. Numerous transformation procedures are known from the literature such as agroinfection using Agrobacterium tumefaciens or its Ti plasmid, electroporation, microinjection or plants cells and protoplasts, microprojectile transformation, to mention but a few. Reference may be made to the literature for full details of the known methods.

Neither is the plant species into which the promoter sequence is inserted particularly germane to the invention. Dicotyledonous and monocotyledonous plants can be transformed. This invention may be applied to any plant for which transformation techniques are, or become, available. The present invention can therefore be used to control gene expression in a variety of genetically modified plants, including field crops such as canola, sunflower,

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tobacco, sugarbeet, and cotton; cereals such as wheat, barley, rice; maize, and sorghum; fruit such as tomatoes, mangoes, peaches, apples, pears, strawberries, bananas and melons; and vegetables such as carrot, lettuce, cabbage and onion. The promoter is also suitable for use in a variety of tissues, including roots, leaves, stems and reproductive tissues.

Thus, nucleic acid sequences which code for novel cysteine proteases have been isolated and characterised. The DNA comprising at least part of the sequence shown in any one of Figures 12 to 17 codes for a cysteine protease and corresponds to the coding region of the sequence. The present invention also includes DNA which shows homology to the sequences of the present invention. The present invention also includes DNA which hybridises to the DNA of the present invention and which codes for at least part of a cysteine protease. Such homology and hybridisation is discussed above in relation to the promoter sequences.

The present invention further includes DNA which is degenerate as a result of the genetic code to the DNA of the present invention and which codes for a cysteine protease.

Also provided by the present invention is a cysteine protease, which is substantially free from other proteins with which it is ordinarily associated, and which is coded for by cysteine protease gene DNA of the present invention.

The present invention will now be described by way of non-limiting example only, and with references to the accompanying drawings, in which:

Figure 1 shows a schematic outline of the identification of cysteine protease isoforms using a reverse transcribed PCR library;

Figure 2 shows the analysis of RT-PCR products by agrose gel electrophoresis and in which DG1 = DEGCYS1 and DG2 = DEGCYS2 oligos;

Figure 3A shows the alignment of the coding regions of the preliminary nucleic acid sequences of RT-PCR clones OSR8.401, OSR8.406, OSR8.403, OSR8.404, OSR8.402, OSR8,389 and OSR8.387;

Figure 3B shows the alignment of the non-coding regions of the preliminary nucleic acid sequences of RT-PCR clones OSR8.389, OSR8.387, OSR8.402 and OSR8.404;

Figure 4 shows the preliminary nucleic acid sequence of clones OSR8.401, OSR8.402 and OSR8.389;

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Figure 5 shows the results of a northern blot of the class 2 clone, OSR8.402, comparing expression in the seed and during germination, using a random primed whole RT-PCR fragment as probe. In this Figure, L = leaf, C = cotyledons, S = seeds and B = buds;

Figure 6 shows the results of northern blots of the class 1 clone, CDCYS12 (using the coding region of the clone as a probe, labelled by PCR), the class 2 clone, CDCYS25 (using the non-coding region of the clone as a probe, labelled by PCR) and the class 6 clone, CDCYS66 (using the non-coding region of the clone as a probe, labelled by PCR) comparing expression in a range of developmental stages. In this Figure, B = bud, W = whole plant and L = leaf;

Figure 7 shows the alignment of the deduced amino acid sequences of clones OSR8.403, OSR8.404, OSR.402, OSR.389, OSR8.387, OSR8.406 and OSR8.401 with the published sequence of COT44 (designated CYS4.BRANA);

Figure 8 shows a schematic outline of the identification of cysteine protease isoforms using a cDNA library;

Figure 9 shows the alignment of the deduced amino acid sequences of the partial length cDNA class 2 clones, CYS2UP6.1, CYS2UP7.1 and CYS2UP8.2, with each other and with COT44 (designated CYS4.BRANA);

Figure 10 shows the alignment of the nucleic acid sequences of the partial cDNA clones, CYS2UP6, CYS2UP7, CYS2UP8, CYS6UP3, CYS6UP5, CYS6UP2 and CYS6UP4, with each other and with COT44;

Figure 11 shows the alignment of the full length cDNA clones, CDCYS66, CDCYS24, CDCYS22, CDCYS25, CDCYS12 and CDCYS14, with each other and with COT44;

Figure 12 shows the nucleic acid sequence of cDNA clone CDCYS12;

Figure 13 shows the nucleic acid sequence of cDNA clone CDCYS14;

Figure 14 shows the nucleic acid sequence of cDNA clone CDCYS22:

Figure 15 shows the nucleic acid sequence of cDNA clone CDCYS24;

Figure 16 shows the nucleic acid sequence of cDNA clone CDCYS25;

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Figure 17 shows the nucleic acid sequence of cDNA clone CDCYS66;

Figure 18 shows the alignment of the predicted amino acid sequences of cDNA clones CDCYS12, CDCYS14, CDCYS22, CDCYS24, CDCYS25 and CDCYS66, and compares the sequences with the characterising features of plant cysteine proteases;

Figure 19 shows a promoter nucleic acid sequence from class 1 genomic clone pKS12p6.

Figure 20 shows a promoter nucleic acid sequence from class 2 genomic clone pKS25p7.

Figure 21 shows a promoter nucleic acid sequence from class 6 genomic clone PKS66P1.

Figure 22 shows the mapping of the transcription start by primer extension experiments as well as the position of the putative TATA box of clones pKS12P6 (class1), pKS25P7 (class2) and pKS66P1 (class 6). Numbers indicate the distance to the 5'end of the cDNAs.

Figure 23 shows the vector constructs for tobacco transformation.

Figure 24 shows the levels of GUS expression in shoot-generating calli during transformation.

Figure 25 shows the levels of GUS expression in leaves from primary transformants.

Figure 26 shows the time course of GUS expression in the progeny of 2 random primary transformants per class, 0 to 36 days after imbibition (DAI). NVS designates the wild type negative control.

Figure 27 shows the GUS activity in the progeny of 24 primary transformants of class 1. Seedlings were assessed at 0, 14 and 28 DAI. The Figure also includes the GUS activity in young leaves from the primary transformants.

Figure 28 shows the GUS activity in the progeny of 24 primary transformants of class 2. Seedlings were assessed at 0, 14 and 28 DAI. The Figure also includes the GUS activity in young leaves from the primary transformants.

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Figure 29 shows the GUS activity in the progeny of 24 primary transformants of class 6. Seedlings were assessed at 0, 14 and 28 DAI. The Figure also includes the GUS activity in young leaves from the primary transformants.

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In general, in the Figures the designations CO or COD in connection with a clone indicate the coding region, and NC or NCD indicates the non-coding region.

In outline, the Examples describe the amplification of a range of germination-expressed cysteine protease partial clones by reverse transcriptase polymerase chain reaction (RT-PCR) on germinating oilseed rape RNA. Preliminary assessment of the clones' expression in dry seeds and germinating seedlings by northern blotting, followed by more detailed northern blot experiments on selected clones to assess their time course of expression. A cDNA library was constructed from tissues showing a high expression of these clones, followed by screening of a genomic library to subclone the promoter areas. Final assessment of the spatial and temporal regulation of the cloned promoters was conducted by transcriptional fusion of the promoter fragments with the  $\beta$ -glucuronidase (GUS) reporter gene and transformation into tobacco.

# Example 1 - CONSTRUCTION OF A CYSTEINE PROTEASE REVERSE TRANSCRIBED PCR LIBRARY FROM GERMINATING OILSEED RAPE SEEDLINGS

In order to identify germination specific sequences, a RT-PCR approach on oilseed rape seedling RNA was utilised. A reverse oligo(dT) primer and forward primers, designed to the cysteine protease coding regions conserved between several plant species, were used to amplify a 750 bp RT-PCR product covering 500 bp of the coding region and about 250 bp of the non-coding region. The 5'-end was sequenced to confirm the identity of the RT-PCR products as cysteine protease related clones. Since there is much less pressure of selection on non-coding regions, significant differences in the 3'-end non-coding regions may predict differences in the 3'-end untranslated regions with an effect on the promoter. This general approach is shown schematically in Figure 1.

### **Preparatory Methods:**

### Plant material

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Five grams of oil-seed rape seeds (*Brassica napus*) from the variety Westar were sterilised in 1% sodium hypochlorite for 10 minutes. After several washes in sterile water, seeds were imbibed with sterile water for 12 hours at 4°C in the dark to synchronise the germination. They were sown on wet sterile Whattman paper and grown at 25°C in the dark prior to harvesting the cotyledons.

### **RNA** extraction

### Lithium chloride method

Total RNA was isolated from dry seeds and three days-old oil seed rape seedlings using a modification of the protocol described by Jepson et al., (1991). Tissues (5 g) were ground with liquid nitrogen in a mortar and pestle until a fine powder was obtained. After addition of 9 ml homogenisation buffer [400 mM NaCl; 50 mM Tris-HCl, pH 9.0; 1 % SDS; 5 mM ethylene diamine tetraacetic acid (EDTA); 4 U/ml heparin; 1 mM aurintricarboxylic acid (ATT); 10 mM dithiothreitol (DTT)] and 4.0 ml phenol saturated in homogenisation buffer and supplemented with 10 % (v/v) m-cresol before use, the tissues were ground again until a fine paste was obtained. The paste was transferred to a cold corex tube and centrifuged for 15 minutes at 13,000 rpm (Sorval, SS34, 4°C). The supernatant was transferred to another tube, extracted for 5 minutes with 5 ml of phenol-chloroform and centrifuged for 30 minutes at 9,000 rpm (Sorval, SS34, 4°C) to recover the aqueous phase. After 3 phenol-chloroform extractions, the RNA was recovered by precipitating the supernatant overnight on ice with one fifth volume of 12 M lithium chloride. After centrifuging for 30 minutes at 9,000 rpm (Sorval, SS34, 4°C), the supernatant was removed and the pellet resuspended in 1 ml of 5 mM Tris (pH 7.5) prior to transfer to a microtube. After a second lithium chloride precipitation overnight, the pellet was washed twice with 70% ethanol, resuspended in 0.2 ml DEPC treated water and stored at -70°C.

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### Caesium chloride method

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Total RNA was isolated from a range of developmental stages of oil seed rape seedlings using a protocol modified from Okayama et al (1979). Tissues (2-4 g) were ground in a mortar and pestle with 1 g Al<sub>2</sub>O<sub>3</sub> in the presence of liquid nitrogen. The powder, kept in dry ice, was then mixed in a corex tube with 8 ml of pre-warmed (65°C) homogenisation buffer [5M thiocyanate guanidine; 0.5 %, w/v, lauryl sarcosine sodium; 0.025 M sodium citrate; pH 7.0] supplemented before use with 2.5 % (v/v) B-mercaptoethanol, and incubated at 40°C for 10 minutes with vortexing until complete defrosting of the tissues. After centrifugation at 12,000 rpm for 30 minutes at 15°C (Sorval, SS34 rotor), the supernatant was recovered, homogenisation buffer added to 8 ml and supplemented with 0.1 g CsCl per ml. After the CsCl had dissolved, the homogenate was added in a 12 ml polyalomer tube containing 2.5 ml of a CsCl high density cushion [5.7 M CsCl; 0.1 M EDTA] without disturbing the cushion. After centrifugation at 25,000 rpm for 24 hours at 20°C (Sorval, TH-641 rotor) the supernatant was removed by suction and the wall of the tube cleaned with absorbent paper before resuspension of the RNA loop into 300 µl resuspension buffer [7 M urea; 2%, w/v, lauryl sarcosin sodium]. The RNA was then transferred to a 1.5 ml tube and extracted with an equal volume of phenol and an equal volume of chloroform/isoamylic alcohol [24:1, v/v]. Following centrifugation at 13,000 rpm for 5 minutes the aqueous phase was recovered and extracted again with an equal volume of chloroform. The RNA was precipitated overnight at -20°C by adding a one-tenth volume of 3 M sodium acetate and 2.5 volumes of cold ethanol. After centrifugation at 13,000 rpm for 15 minutes at 4°C, the supernatant was discarded and the pellet was washed with 1.5 ml of cold 70% ethanol. The centrifugation was repeated for 10 minutes, the supernatant was discarded and the pellet dried for 3-4 minutes in a speedvacuum. The pellet was resuspended in sterile DEPC-treated water, the RNA precipitated again and the pellet washed, as described above, and finally resuspended in 50 µl DEPC-treated water and stored at -70°C.

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### Labelling of DNA probes

### Terminal exchange

The oligos (25-50 ng) were labelled by phosphorylating their hydroxylated 5'-end using 20 U T4 polynucleotide kinase (New England Biolabs) in a 25 μl reaction [30 uCi [γ
32P] ATP (Amersham, 5000 Ci/mmol), 1X kinasing buffer] at 37°C for 30 minutes. The oligos were purified from the unincorporated nucleotides using G-25 sephadex spun columns (5prime→3prime, Inc®), as recommended by the manufacturer.

### **PCR**

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Plasmid DNA (5 ng) was amplified [(94°C, 1 minute; 65°C, 1 minute; 72°C, 1 minute) x 17 cycles; (72°C, 7 minutes) x 1 cycle] using 2.5 U of Taq polymerase (Gibco BRL) in a 50 μl PCR reaction [0.25 μM [α-32P] dATP(Amersham, 3000 Ci/mmol); 0.4 μM dATP; 50 μM other dNTPs; 1.5 mM MgCl2; 0.5 μM oligos; 1X PCR buffer]. The probes were purified from the unincorporated nucleotides using G-50 sephadex spun columns (Pharmacia Biotech), as recommended by the manufacturer.

### 15 Random priming

Plasmid DNA (25-50 ng) was labelled using the "Oligolabelling Kit" (Pharmacia) following manufacturer's recommendations.

### Northern blot hybridization

Northern blot experiments were performed according to Sambrook *et al.* Total RNA (10 μg) was mixed with 2.5 vol of loading buffer (5prime→3prime, Inc®), sized by electrophoresis on a 1.2 % agarose denaturing gel [1X MOPS(40 mM MOPS, pH 7.0; 10 mM sodium acetate; 1 mM EDTA); 17%, v/v, formaldehyde] in alkaline running buffer [1X MOPS; 7%, v/v, formaldehyde] and transferred to nylon membranes (Hybond N, Amersham) by capillary blotting according to the manufacturer's recommendations. Hybridizations were performed overnight at 65°C in hybridisation buffer [5XSSPE (900 mM NaCl; 50 mM NaH<sub>2</sub>PO<sub>4</sub>.H<sub>2</sub>O; 5 mM EDTA; pH 7.4, NaOH); 0.5% sodium dodecyl sulphate (SDS); 1% powdered milk] supplemented with 100 μg/ml denatured salmon sperm DNA. Blots were

washed in 3X SSC [20X SSC: 450 mM NaCl; 45 mM Na<sub>3</sub>citrate.2H<sub>2</sub>O; pH 7.0, HCl] and 1% SDS for 1 hour at 65°C, and exposed to X-ray films (X-OMAT AR, Kodak) at -80°C with intensifying screen.

### **RT-PCR Library Construction and Screening**

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A RT-PCR library was constructed from germinating oilseed rape seedlings using a "Perkin Elmer GeneAmp® RNA PCR Kit" and following the manufacturer's recommendations. Total RNA (1 μg) extracted from cotyledons 3 days after seeds imbibition was reverse transcribed [65°C, 2 minutes; 42°C, 30 minutes; 99°C, 5 minutes; 6°C, 5 minutes] using an oligo(dT) primer (MPRACE1B) stabilised for the PCR. The heteroduplex DNA-RNA was subsequently amplified by PCR using the same downstream primer as for the reverse transcription and a set of two upstream degenerate primers (DEGCYS1 and DEGCYS2) designed on the peptide level from a motif of cysteine protease coding regions conserved among most of the plant species [GCNCCLM(NED)]. Cycling conditions: [(95°C, 2 minutes; 55°C, 2 minutes; 72°C, 1.5 minutes) x 2 cycles; (95°C, 2 minutes; 55°C, 1 minute; 72°C, 1.5 minutes) x 33 cycles] followed by 7 minutes at 72°C.

The RT-PCR products were ligated into a pCRII vector before transforming *E. coli* using the "TA Cloning® Kit" (Invitrogen) following the manufacturer's recommendations. This system takes advantage of the non-template dependent activity of thermostable polymerases used in PCR that add a single deoxyadenosine to the 3'-ends of all duplex molecules provided by PCR. This allows direct cloning into a pCRII vector which contains overhanging deoxythimidine. Plasmid DNA was purified using "Wizard DNA miniprep DNA purification system" (Promega) and sequenced by the chain termination method (Sanger *et al.*, 1977) with "Sequenase version 2.0 T7 DNA polymerase" (USB) according to the manufacturer's recommendations.

OLIGOS	SEQUENCE $(5' \rightarrow 3')$
MPRACE1B	GGC CAC GCG TCG ACT AGT TAC TCG AGT
	TIT TIT TIT TITTIT T
DEGCYS1	GGI TG(CT) AA(CT) GGI GGI (CT)TI ATG
DEGCYS2	GGI TG(CT) AA(CT) GGI GGI (CT)TI ATG (GA)A

### Analysis

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The RT-PCR products were analysed by electrophoresis on an agarose gel. As can be seen in Figure 2, two major 750 bp fragments of the expected size, amplified from cotyledons RNA, but not from seeds RNA were found. These were gel excised and cloned. In a parallel approach, aliquots of the RT-PCR products without gel purification were cloned to generate a three days-old expressed cysteine protease library containing 400 clones from which 22 came from gel excision. A colony screening with the oligos used for the RT-PCR identified 250 putative cysteine proteases. Seven clones from gel excision were taken at random and fully sequenced, all of them were cysteine proteases, and fell into 3 classes.

These clones were given the following designations:

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CLONES	CLASS
OSR8.401	1
OSR8.402	2
OSR8.403	2
OSR8.404	2
OSR8.406	6
OSR.387	6
OSR.389	6

The preliminary DNA sequences of these clones and their alignment is shown in Figures 3A and 3B.

Three clones, one per class, OSR8.401 (class 1); OSR8.402 (class 2) and OSR8.389 (class 6) whose preliminary DNA sequences are shown in Figure 4, were labelled by random priming and assessed on northern blots containing total RNA extracted from seeds and cotyledons using the lithium chloride method. They appeared to be well expressed during germination but not in dry seeds. Furthermore, some cross-hybridisation was observed

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between the different classes due to the nature of the probe. This is illustrated in Figure 5 which shows the result for the class 2 clone, OSR8.402, with a random primed whole RT-PCR fragment as probe.

The library was then screened using the five cysteine protease clones labelled by random priming. No new classes were clearly identified, all the diversity of clones was present in the gel-excised fragments. The expression pattern of the three classes of clones was assessed by northern blots with RNA, extracted with the caesium chloride method, from a range of developmental stages as shown in Figure 6. For class 2 and 6, hybridisations were performed using the 3'-non coding regions of clones OSR8.402 and OSR8.389 respectively, to avoid any cross-hybridisation. For class 1, a part of the coding region of OSR8.401 clone was used as a probe because no non-coding region was available in the RT-PCR clones. To increase the specificity, probes were labelled by PCR using the oligo(dT) reverse primer (MPRACE1B) and an internal reverse primer (CYS8.402 and CYS8.389) for classes 2 and 6, whilst using 2 internal primers (CYS8.401 and CYS8.401R) for class 1. Classes 2 and 6 are expressed following seed imbibition and for the first 4-5 days of early seedling growth but are not expressed in mature plant organs or in the developing seed. Class 1 shows some expression in buds and leaves but this may be the result of some cross-hybridisation due to the nature of the probe. Class 2 is highly related to COT44, the only cysteine protease published for oilseed rape (Comai and Harrada, 1989). Figure 7 shows alignment of deduced amino acid sequences of the clones with COT44.

OLIGOS	SEQUENCE $(5' \rightarrow 3')$
CYS8.401	TAT CCT TAT CAA GAA CGT GAT GGC A
CYS8.401R	CCT ACG ATG AGC ACT GCG TGG T
CYS8.402	GCA GTA ATC AAA TTG GGA TTG TTA TAA
CYS8.389	CGT GGA ACC AGC AGT GTT TGA AGT T

# Example 2 - CONSTRUCTION OF A STANDARD OLIGOd(T) PRIMED AND A CYSTEINE PROTEASE SPECIFICALLY PRIMED cDNA LIBRARY FROM GERMINATING OILSEED RAPE SEEDLINGS

As the RT-PCR products were not full length and to avoid PCR generated mutations, an oilseed rape cDNA library was constructed from a developmental stage showing a high expression of the three clones of interest. A specifically primed library was constructed using two specific oligos designed on the basis of the three classes of RT-PCR clones, rather than using an oligo(dT) primer. As a result only the cysteine protease clones were reverse transcribed, which provided a small number of short clones of about 650 bp, all of them being full length at the 5'-end. This allowed the design of oligos to the 5'-non-coding regions for use in screening a standard oligo(dT) primed cDNA library directly for full length clones. This general approach is shown in schematically in Figure 8.

### **Preparatory Methods:**

### 15 Plant material

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Five grams of oilseed rape seeds (*Brassica napus*) from the variety Westar were sterilised in 1% sodium hypochlorite for 10 minutes. After several washes in sterile water, seeds were imbibed with sterile water for 12 hours at 4°C in the dark to synchronise the germination. They were sown on wet sterile Whattman paper and grown at 25°C in the dark for 2 days prior to harvesting the cotyledons.

### RNA extraction and purification

Total RNA was isolated from 2 days-old oil seed rape seedlings using the caesium chloride method described previously. Polyadenylated RNA was purified from 1 mg of total RNA using a "PolyATract mRNA isolation system" (Promega), according to the manufacturer's recommendations. The system uses a biotinylated oligo(dT) primer to hybridise at high efficiency in solution to the 3'-poly(A) region of the mRNAs. The hybrids were then captured and washed at high stringency using streptavidin coupled to paramagnetic particles and a magnetic separation strand, prior to elution with water.

### cDNA Library Construction and Screening

The standard oligo(dT) primed cDNA library and the cysteine protease specifically primed library were constructed from 2 days-old oilseed rape poly(A) RNA (5 µg), using a lambda "ZAP-cDNA® Synthesis Kit" (Stratagene). The manufacturer's recommendation were followed strictly although, for the specific library, the reverse transcription was primed using a mix of two specific oligos (CDNA8.401R and CDNA8.387R) respectively for the class 1 and for classes 2 and 6 of the RT-PCR clones, modified to include a *Xho* I site at their 5'-end. The second strand was synthesised by nick-translation using DNA polymerase I, after treatment of the heteroduplex with RNase H. The cDNAs were filled in with Klenow, ligated to *EcoR* I adapters and digested with *Xho* I prior to size-fractionation on Sepharose®-400 spun column (Pharmacia) and directional cloning as an *EcoR* I-Xho I insert into the polylinker of pBluescript phagemid contained within the Uni-ZAP vector arms. Lambda-ZAP is a replacement lambda which has been engineered to contain pBluescript phagemid, which polylinker is used for cloning the cDNAs.

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OLIGOS	SEQUENCE $(5' \rightarrow 3')$
CDNA8.401R	GAG AGA GAG AGA GAA CTA GTC
	TCG AGT CCC ATG GTT TTT AAT
CDNA8.387R	GAG AGA GAG AGA GAA CTA GTC
	TCG AGC CGC CGT TTT TCA T

The library was packaged *in vitro* using Gigapack® II Gold packaging extract (Stratagene), plated on *E. coli* cell line XL1-Blue MRF' and transferred onto nylon membranes (Hybond N, Amersham) according to the manufacturer's recommendations. Labelling of probes, hybridisations and washes were performed as described previously.

Selected lambda-ZAP clones were excised *in vivo* to recover the cloned cDNA as a phagemid in pBluescript SK. E. coli SolR cells were co-transfected with the recombinant lambda-ZAP and with a helper phage which provided the proteins necessary for the synthesis of a single strand of DNA which, once circularised, provide a functional phagemid.

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### Analysis

a) A small cysteine protease-enriched specifically primed cDNA library containing 1.10<sup>4</sup> plaque forming units (pfu) was obtained from 2 days-old oilseed rape cotyledons. 5.10<sup>3</sup> pfu were plated and transferred onto three replicate membranes to check for cross-hybridisation. Membranes were screened with 3 oligos (CYS8.401MR, CYS8.402MR, CYS8.406MR) designed respectively to the 5'-end of classes 1, 2 and 6 of the RT-PCR clones and labelled as described above.

Five similar but distinct 5'-end cysteine protease cDNAs containing a short 5'-untranslated region and 650 bp of coding region were obtained, excised *in vivo* and sequenced. They fall into class 2 and class 6 but no 5'-end cDNA was found for class 1. Although clones from class 2 are highly related to COT44 cDNA (Comai and Harrada, 1989), their 5'-end is 160 bp longer. This indicates that COT44 is missing, 46 amino acids (aa) corresponding to the signal peptide and to a part of the propeptide. This is further illustrated by Figure 9 which shows the alignment of the deduced amino acid sequence of the cDNA clones CYS2UP6, CYS2UP7 and CYS2UP8 from class 2 with COT44.

Two oligos (CYS6B-UP and CYS6A-UP) were designed to the 5'-non-coding region of respectively, cDNAs from classes 2 and 6, to screen the standard oligo(dT) primed library directly for full length clones. The preliminary sequence alignment of the clones with each other and COT 44 is shown in Figure 10.

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OLIGOS	SEQUENCE $(5' \rightarrow 3')$		
CYS8.401MR	CCC ATG GTT TIT AAT GAC AAA TTG AAA A		
CYS8.402MR	CCG CCG TTT TTC ATT ATG AAT TGA AA		
CYS8.406MR	CGC CGT TTT TCA TGA TGA ATT GAA AA		

b) A representative oligo(dT) primed cDNA library containing 1.10<sup>7</sup> pfu was obtained from 2 days-old oilseed rape cotyledons. The size of the inserts, estimated by PCR, is ranging from 0.75 kb to 3 kb with an average insert size of 1.5 kb. 1.10<sup>6</sup> pfu were plated and transferred onto three replicate membranes to check for cross-hybridisation. Membranes were first screened with a 5'-end oligo (CYS8.401MR, CYS6B-UP and CYS6A-UP) designed respectively to the 5'-end coding region of class 1 RT-PCR clones and to the 5'-

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untranslated region of class 2 and class 6 cDNAs to identify full length clones for class 2 and 6 (class 1, 20 positives; class 2, 250 positives; class 6, 130 positives).

OLIGOS	SEQUENCE $(5' \rightarrow 3')$
CYS8.401MR	CCC ATG GTT TTT AAT GAC AAA TTG AAA A
CYS6B-UP	TAG AAA ACC AAC AAA ACA AAC ATA CAA T
CYS6A-UP	GAA CAA CCA AGC CAA ACA TAC AAT AT

The same membranes were screened again with the 3'-end probes used on the developmental northern blots to ensure the correct form was chosen. For classes 2 and 6, hybridisations were performed using the 3'-non-coding regions labelled by PCR to avoid any cross hybridisation. For class 1, a part of the coding region was used as a probe because non-coding region was not available in the RT-PCR clone (class 1, 84 positives; class 2, 205 strong positives; class 6, 85 strong positives).

Comparison between the number of clones identified with both specific probes (5' and 3') confirmed that most of the cysteine protease cDNA clones present in the library are full length.

Ten clones per class which hybridised with (5' and 3' probes) but did not cross hybridise with probes from the two other classes were plaque purified and four clones per class were excised *in vivo*. Six full length cysteine protease cDNA clones falling into the three classes of cysteine proteases identified from the RT-PCR work were isolated and fully sequenced (two for class 1, three for class 2, and one for class 6). The alignment of the cDNA clones is shown in Figure 11. They fall into 3 classes of CP related to the papain super family and the pre-proenzymes share 52% (class 1), 90% (class 6) and 96% identity (class 2) with cot44. Figures 12 to 17 show the nucleic acid sequences of clones CDCYS12, CDCYS14, CDCYS22, CDCYS24, CDCYS25 and CDCYS66 respectively. The peptide sequences were predicted and showed the characteristic features present in most of the plant cysteine proteases, as shown in Figure 18 for clones CDCYS12, CDCYS14, CDCYS25, CDCYS25 and CDCYS66.

# Example 3 - SCREENING OF AN OILSEED RAPE GENOMIC LIBRARY AND SUBCLONING AND CHARACTERISATION OF THE PROMOTER REGIONS

Oligonucleotide probes were generated to the 5'-end non-coding region of one cDNA clone per class and used to screen a genomic library in order to isolate clones carrying the promoter regions. For each class, genomic clones were isolated and the promoter subcloned into a phagemid for more precise characterisation and deletion.

### Genomic Library Construction and Screening

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An amplified  $\lambda$ EMBL-3 random genomic library (Clontech) from oilseed rape (Brassica napus cv. Bridger) was constructed. DNA was partially digested with Mbo I and the fragments were separated on a sucrose gradient to produce size range between 8 to 22 kb before cloning into the BamH I site of a  $\lambda$ EMBL-3 replacement vector. The library was plated on E. coli strain LE392 cells. Screening and plaque purification were performed as described by Sambrook et al. (1989). Genomic clones corresponding to the three classes of cDNAs were isolated and  $\lambda$ DNA minipreparations were carried out using a protocol from Grossberger (1987). Genomic clones were mapped using their restriction fragment length polymorphism (RFLP) patterns: clones were digested with a set of restriction enzymes, analysed on a 0.8% agarose gel and simultaneously transferred onto two membranes (Hybond N, Amersham) according to Sambrook et al., (1989) prior to hybridisation.

### Analysis

For the primary screening twenty genome-equivalent (2.10<sup>6</sup> pfu) were plated and transferred onto three replicate membranes to check for cross-hybridisation. Membranes were hybridised with 3 oligos (CDNA12, CDNA25 and CDNA66) designed respectively to the 5'-end of classes 1, 2 and 6 of the cDNA clones, to get as close as possible to the promoter area (class 1, sixteen strong positives; class 2, nine strong positives; class 6, eight strong positives). No cross-hybridisation was detected between the three classes and ten clones per class were chosen for a secondary screening.

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OLIGOS	SEQUENCE $(5' \rightarrow 3')$
CDNA12	ATC GTC TTC CTT TGT TTC TCT CA
CDNA25	CTT CGT CAG CGA AAC TCC TCT CTT
CDNA66	CAG AAC TAG AAC AAC CAA GCC AAA C

The secondary screening was performed using 3'-end PCR probes as described in connection with Figure 6, to specifically detect the genomic clones corresponding to the RT-PCR clones assessed by northern blots. Some of the clones were not identified by these probes (one for class 1 and two for class 6), since these were likely to be short clones in their 3'-end, thus useful for promoter isolation, they were rescued using the probes from the primary screening. Ten clones per class were plaque purified by two more rounds of purification using the probes from the primary screening. In order to avoid redundant clones (amplified, random library), DNA was prepared from eight genomic clones per class and characterised by RFLP. Clones were cut with *Sal* I and *Bam*H I, analysed on a 0.8% agarose gel, transferred onto replicate membranes and hybridised with two sets of probes. Oligos were designed to the 5'-non-coding region (CDNA12, CYS6B-UP and CYS6A-UP) and to the middle of the coding region (CYS8.401MR, CYS8.406MR and CYS8.406MR) of cDNA clones CDCYS12, CDCYS25 and CDCYS66 respectively.

Four remaining clones per class (12g4, 12g5, 12g6, 12g8; 25g2, 25g4, 25g5, 25g7; 66g1, 66g4, 66g8 and 66g9) were further characterised using another round of digestion/hybridisation. Class 1 cDNAs contain a *Bgl* II site 500 bp from the translation start and class 2 and 3 cDNAs contain a *Hind* III site 300 bp from the translation start, these enzymes were used in association with and without *Sal* I, which releases the insert, to generate genomic fragments suitable for subcloning. PCR experiments were carried out on genomic DNA and cDNAs to predict the size of the promoter area by identifying putative introns. PCR, using a forward primer in the 5'-non-coding region and a reverse primer located after the *Bgl* II and *Hind* III restriction sites, showed the presence of a 400 bp intron within the first 600 bp of class 1 cDNAs whilst no intron was present within the first 300 bp of class 2 and class 6. Promoter fragments with a predicted size in the range of 2-5 kb were identified for one genomic clone per class (12g6, 25g7 and 66g1), ready to be subcloned into pBluescript KS<sup>+</sup>.

## Example 4 - CHARACTERISATION OF TRANSCRIPTION STARTS

Genomic lambda-fragments containing the promoter were subcloned into pBluescript KS<sup>+</sup> for more precise characterisation. Sequencing allowed the identification of putative transcription signals before mapping the actual transcription start by primer extension experiments. This involved the extension of a labelled reverse primer designed in an area close to the translation start. After degradation of the RNA template the extension products were sized in a polyacrylamide gel.

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### **Analysis**

Genomic fragments containing the promoter were subcloned into pBluescript as a Bgl II-2.6 kb insert cloned in BamH I for class 1 (pKS12P6), as a Hind III-4.2 kb insert for class 2 (pKS25P7) and as a BamH I-Hind III-2.4 kb insert for class 6 (pKS66P1). Sequencing with pUC1 and pUC4 vector oligos and with two internal reverse primers designed to the 5'-end of the cDNAs (CDNA14R for class 1 and CDNA66R for class 2 and 6), allowed the orientation of the clones and the identification of putative transcription signals (Pautot et al.,1989). The full nucleotide sequence of the promoters from the sub-cloned genomic fragments is given in Figures 19, 20 and 21 respectively for class 1, 2 and 6.

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OLIGOS	SEQUENCE $(5' \rightarrow 3')$
CDNA14R	GAA GAA ACT AGA AGA AGG GAG AAG AA
CDNA66R	TCA CTT CTT CAT CGG TTC TCC A

Transcription starts have been mapped precisely by primer extension experiments according to Sambrook *et al.* (1989) modified as follows. Oligos (CYSGE12R, CYSGE25R and CYSGE66R) designed respectively for class 1, 2 and 6 were labelled by terminal exchange as described previously. Total RNA (50 ug), isolated from 3 days-old oilseed rape cotyledons using the caesium chloride method, was precipitated together with 2 ng of primers and resuspended in 30 µl hybridisation buffer [1 mM EDTA; 400 mM NaCl; 40 mM Pipes, pH 6.4; 70%, v/v, deionised formamide]. Annealing was performed overnight at 32°C

following denaturing at 85°C for 10 minutes. After precipitation and resuspension in 25 μl of reverse transcription buffer [50 mM KCl; 10 mM MgCl<sub>2</sub>; 1 mM dNTPs; 1 U/ul RNase inhibitor], the primers were extended for 90 minutes at 42°C with 2.5 U MuLV reverse transcriptase (Perkin-Elmer). Template RNA was degraded for 30 minutes at 37°C with 20 U of RNase (RNace-it, Stratagene). For each class, the extension products were analysed on a polyacrylamide denaturing gel in parallel with a sequencing reaction performed on the genomic clones (pKS12P6, pKS25P7 and pKS66P1) using the same primers as for the primer extension.

OLIGOS SEQUENCE $(5' \rightarrow 3')$		
CYSGE12R AGG AAG AAG ACG ATG ATG GTG ACA		
CYSGE25R	GTA CAA GAG AAG TAA AGA GAG GAG T	
CYSGE66R	CGT ATA GGA GAA GTA AAG AAA TGA GT	

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As shown in Figure 22, class 1 transcription start is in a good context [t<sub>27</sub>T<sub>35</sub>C<sub>49</sub>A<sub>78</sub>a<sub>18</sub>C<sub>45</sub>g<sub>8%</sub>] compared to 49 other plant genes compiled by Pautot *et al.*(1989). Conserved nucleotides are in capitals, important ones are in bold and the transcription start point is underlined. This transcription start has been mapped 22 nucleotides downstream a putative TATA box localised 179 nucleotides before the translation start and 152 nucleotides upstream the longest cDNA.

Although the consensus for the TATA box is not optimal [C<sub>34</sub>a<sub>18</sub>t<sub>32</sub>t<sub>34</sub>a<sub>3</sub>A<sub>97</sub>T<sub>90</sub>A<sub>94</sub>a<sub>47</sub>A<sub>95</sub>a<sub>30</sub>A<sub>71</sub>G<sub>44%</sub>] compared to 79 other plant genes compiled by Pautot *et al.* (1989), this result is a confirmation of a previous primer extension experiment using an oligo priming 34 nucleotides downstream the ATG. The distance between the transcription start and the longest cDNA might be explained by the presence of an intron within the 5'untranslated region or by the existence of an alternative transcription start point.

The Class 2 transcription start is in a good context  $[a_{18}a_{20}B_{22}A_{78}T_{49}C_{45}A_{43\%}]$ , and has been localised 33 nucleotides after a putative TATA box fitting very well within the plant consensus  $[T_{37}g_{11}g_{14}t_{34}T_{96}A_{97}T_{90}A_{94}a_{47}A_{95}T_{63}A_{71}G_{44\%}]$ . This corresponds to 53 nucleotides before the translation start and 26 nucleotides upstream the cDNA (Figure 22).

Class 6 transcription start is nearly in the same context as class 2

[g<sub>18</sub>a<sub>20</sub>a<sub>22</sub>A<sub>78</sub>T<sub>49</sub>C<sub>45</sub>A<sub>43%</sub>] and has been localised 30 nucleotides after a putative TATA box

showing exactly the same consensus as for class 2. This corresponds to 51 nucleotides before the translation start and 19 nucleotides upstream the cDNA (Figure 22).

# **Example 5 - PROMOTER EXCISION**

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Prior to fusion with the reporter genes, promoters must be cut precisely between the transcription start and the translation start. Since no useful restriction site was available for the class 2 and 6 genomic clones, a site was engineered into a PCR fragment used to replace a corresponding endogenous fragment.

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### **Analysis**

A Hind III site was introduced by PCR on class 1 genomic clone, 2 nucleotides before the translation start, in order to eliminate the remaining part of coding region. The 270 bp fragment was generated by 15 cycles of PCR on pKS12p6 DNA, using CYSGE12C and CYSG12CR oligos.

In the same way, a *BamH* I site was introduced before the translation start of class 2 and class 6 genomic clones using 2 sets of oligos (CYSGE25C, CYSG25CR and CYSGE66C, CYSG66CR) respectively on pKS25p7 and pKS66p1 DNA to generate a 165 bp fragment.

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For class 1, pKS12p6 was cut with Bsm I and Hind III and gel recovered to excise a 900 bp fragment prior to replacement with the PCR fragment cut by Bsm I and Hind III, to generate pKS12P.

For class 2, pKS25p7 was cut with Sph I and BamH I, gel recovered to excise a 460 bp fragment and ligated to the replacement PCR fragment cut by Sph I and BamH I, to generate pKS25P.

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For class 6, pKS66p1 was cut with *Hind* III, filled in with Klenow fragment of DNA polymerase I, cut with *Sph* I and gel recovered to excise a 440 bpfragment. The PCR fragment was blunt ended with T4 DNA polymerase, cut with *Sph* I and cloned into the deleted pKS66p1 to generate pKS66P.

OLIGOS	SEQUENCE $(5' \rightarrow 3')$
CYSGE12C	GTA ATG GCC TAG CCT GTC TGG C
CYSG12CR	GAT GAT GGT GAC AAG CTT TTT CTT ACA GG
CYSGE25C	CTA TCT TGC ATG CCC ATT ATT ACT TT
CYSG25CR	ACG AAG CCG GAT CCT ATG TTT GTT TTG TTG
CYSGE66C	CAT CTT GCA TGC CCA TTA CTG CAT
CYSG66CR	AGG AAG CCG GAT CCT ATG TTT GGC TTG G

### **Example 6 - PROMOTER-GUS CONSTRUCTS**

In order to assess the spatial and temporal regulation of the cloned promoter regions in an heterologous system and under different biotic and environmental conditions, they were used them to drive a reporter gene into tobacco. Transcriptional fusions between each promoter fragment and the  $\beta$ -glucuronidase (GUS) gene were engineered for plant assays and histochemical localisation. For the avoidance of doubt, a reporter gene is used here for convenience only, and to demonstrate the principles involved. In non-test situations the gene controlled by the promoter of the present invention will be that which produces the desired effect.

### **Plasmid Construction**

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Standard recombinant DNA methods were adopted in the construction of plasmid vectors (Sambrook et al., 1989). The CP12 Hind III-Not I-1.7 kb promoter fragment was excised from pKS12P, filled in using Klenow fragment of DNA polymerase I and ligated into the Sma I site of the Agrobacterium Ti vector pTAK1 containing the E.coli uidA gene encoding β-glucuronidase (Jefferson et al., 1987), to produce a pTAKCP12 binary vector. In the same way CP25 Hind III-BamH I-3.7 kb and CP66 BamH I-1.9 kb promoter fragments were excised from pKS25P and pKS66P respectively, and ligated into pTAK1 cut with the same enzymes to produce pTAKCP25 and pTAKCP66 binary vectors. All constructs were transformed into E. Coli strain DH5α as an intermediate host for the vectors construction. The structure of the resultant chimeric reporter gene constructs was verified by PCR, restriction digest and sequence analysis. Figure 23 shows a schematic of the constructs for plant transformation.

### **Plant Transformation**

Plasmids pTAKCP12, pTAKCP25 and pTAKCP66 were transferred into Agrobacteriun tumefaciens LBA4404 using the freeze/thaw method described by Holsters et al. (1978). Tobacco (Nicotiana tabacum var. Samsun) transformants were produced by the leaf disc method (Bevan, 1984). Shoots were regenerated on medium containing 100mg/l kanamycin and 200 mg/l carbenicillin. After rooting, plantlets were transferred to the glasshouse and grown under 16 h light/8 h dark conditions.

### Results

	12CP plants	25CP plants	66CP plants
# shoot taken	440	250	404
# shoot rooting	130 (30%)	82 ( 33%)	120 (30%)
# shoot subdivided	100	82	100
# shoot re-rooting	84 (84%)	58 (70%)	71 (71%)

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### PRIMARY TRANSFORMANT ANALYSIS

### **Objective**

Primary transformants were first analysed by polymerase chain reaction (PCR) in order to reduce the number of plants to analyse and to make sure they contained the intact promoter-reporter gene cassette.

Since promoters can be deregulated in callus, GUS analysis were carried out on calli from transformation to make sure the promoters are not active at this stageas this might have had a deleterious effect on future transformations efficiency depending on the nature of the transgene (e.g. if barnase ribonuclease gene is driven by one of these promoters). Promoter activity was also assessed in young leaves from primary transformants to confirm the absence of ectopic expression at this stage.

### Polymerase chain reaction

Genomic DNA for PCR analysis of transgenic plants was prepared according to Edwards et al. (1992). Plant extracts DNA (2.5 ul) was amplified [hot start at 80°C; (94°C, 1

min; 63°C, 1 min; 72°C, 1 min) x 35 cycles; (72°C, 7 min) x 1 cycle] using 2 U of Taq polymerase (Gibco BRL) in a 25 ul PCR reaction [200 uM dNTPs; 3 mM MgCl2; 1 uM oligos; 1X PCR buffer].

### 5 Results

A total of 37 individual transformants per class were randomly picked from *in-vitro* culture 12CP, 25CP and 66CP explants and analysed with 2 sets of primers. The first set contained one primer specific to the 5' end of the NOS terminator of the NPTII gene (NOSTER1) and a reverse-primer specific to the 5'end of the cloned promoters (CYSGE12R, CYSGE25R or CYSGE66R). The second set contained one primer specific to the 3'-end of the promoters (CYSGE12C, CYSGE25C or CYSGE66C) and a reverse-primer specific to the 5' portion of the GUS gene (GUS1R). A total of 34 explants was found to be double PCR positive for class 6 (94% of the plants tested), while for class 1 and class 2 only 27 plants gave the expected result (73% of the plants tested). Plants containing the intact cassette were transferred to the glasshouse and self-pollinated.

OLIGOS	SEQUENCE ( $5' \rightarrow 3'$ )
CYSGE12RT	GGG TTC TTC TGG GTA GCA AAC TG
CYSGE25RT	ACT TCA CGT TCT GAA TCT CAT CGA A
CYSGE66RT	GGG CCA GAA TGC GGA TTT TAC TAA
GUSIR	CGC TTT CCC ACC AAC GCT GAT C
NOSTER 1	TTG AAT CCT GTT GCC GGT CTT GC

### GUS enzyme assays

Fluorometric assays for GUS activity performed with the substrate 4-methylumbelliferyl-D-glucuronide (Sigma) were carried out using a Perkin-Elmer LS-35 fluorometer (Jefferson *et al.*, 1987). Protein concentration of tissue homogenates were determined by the Bio-Rad protein assay (Bradford, 1976).

### Results

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GUS assays were carried out for each class on 20 regenerating calli resulting from the transformation process. Tobacco extracts from wild type leaves and callus as well as from leaves from 35S-GUS transgenic plants were used respectively as negative and positive

controls. As shown in Figure 24, no significant GUS activity could be detected in calli compared to the levels presents in leaves from the 35S-GUS plants.

Figure 25 shows a preliminary assessment of the levels of GUS activity in young leaves from primary transformants of each class. Results indicate that the promoters are not active at this stage. This is a confirmation of the northern blot results obtained for class 2 and 6 but contradicts those for class 1 (Figure 6). Class 1 CP showed some expression in leaves but it is thought to be due to a cross-hybridisation problem due to the nature of the probe. The GUS result seemed to confirm the latest hypothesis.

### **ANALYSIS OF SEGREGATING POPULATIONS**

### Objective

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The objective was to select for each class, four GUS expressing lines ranging from low expressors to high expressors, preferably from lines with a single locus insertion of the transgene as this facilitates the comparisons between lines. The number of loci in the primary transformants is estimated by the segregation of the NPTII (kanamycin-resistance) gene in the progeny.

### Segregation test

Seeds were sterilised in 10% bleach for 15 min. After several washes in sterile water around 150 seeds were sown on ½ MS medium (2.3 g/l MS salt, 1.5% sucrose, 0.8% Bactoagar, pH 5.9) containing 100 mg/l kanamycin. Seeds were grown for three weeks at 26°C with 16 hours/8 hours light/dark prior to scoring. If the primary transformants contained one copy of the transgene, the expected ratio for kan<sup>R</sup> to kan<sup>S</sup> seeds was 3 to 1 (although in very rare cases one locus could possibly contain several transgenes).

### 25 Results

A substantial number of plants are showing a *petaloïdie* phenotype in which a variable proportion of the flowers in a plant have one or more normal stamens replaced by petals.

Some of these plants were so badly affected that we could not recover any seeds.

The table below summarises the genetic data for the primary transformants.

	12CP plants	25CP plants	66CP plants
# lines in glasshouse	27	27	34
# lines giving seeds	22 (81%)	22 (81%)	29 (85%)
# petaloïdie phenotype	13 (48%)	11 (41%)	5 (15%)
# single loci insertion	11 (50%)	9 (41%)	12 (41%)

## Preliminary time course experiment

The northern blot results indicated an accumulation of CP mRNA in oil seed rape at 2 to 3 days after seeds imbibition (DAI). However, the heterologous expression in tobacco may differ from the endogenous expression in oil seed rape due to differences in physiology and transcription machinery. Furthermore, the activity of the promoter was indirectly analysed through a reporter protein, which delays the detection. So, in order to work out at which time point all the F1 generations should be analysed, the time course of GUS expression from each promoter had to be established.

#### Results

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The experiment was carried out on 2 random lines per class as well as on wild type and 35S-GUS control lines. For each time point, 40 seeds of lines 12CP5, 12CP14, 25CP8, 25CP13, 66CP8 and 66CP76, as well as the controls, were grown at 26°C with 16 hours/8 hours light/dark, on plates containing 1/2 MS media. Seedlings were sampled at 0, 2, 4, 6, 8, 10, 29 and 36 DAI and stored at -70°C. At time point 0 to 10 DAI, the 10 biggest seedlings were collected while only 5 were taken at 29 and 36 DAI which increased the variability within these samples.

In tobacco seedlings grown under the conditions described above, three lines out of 6 showed an induction of GUS expression during germination (Figure 26). The activity peaked at around 30 DAI, although the protein accumulation clearly started at 8 DAI. Thus, the F1 generations should be assessed at 14 and 28 DAI to identify a range of GUS expressors. The maximum of activity was about 5% of 35S-GUS in the best expressing line (12CP5) although the relatively high level of expression in leaves suggests that this might be due to a position

effect of the transgene. Normal levels of expression are more likely to be around 1% of 35S-GUS.

# Identification of high GUS-expressing lines during germination.

### 5 Results

Seedlings were grown on 1/2 MS media supplemented with 100 mg/l kanamycin in the conditions described previously. Five seedlings were harvested at 0 (dry seeds), 14 (2 expanded leaves) and 28 DAI (4 expanded leaves), pooled and assessed in duplicates as described previously.

Figures 27, 28, and 29 summarise the expression levels for class 1, 2 and 6 respectively. These preliminary data suggest that the promoters are expressed in a seedling-specific manner in tobacco. As expected from the RNA study in oilseed rape the levels of expression are low. Class 2 promoter fragment is more active than class 1 at this stage, while class 6 gives extremely low levels of expression.

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### **GUS** histochemical detection

GUS histochemical staining of whole seedlings was achieved by vacuum infiltration with a solution containing 1 mM X-gluc (5-bromo-4-chloro-3-indolyl -D-glucuronide, 100 mM NaPO4 pH 7.5, 10 mM EDTA, 0.5 mM K3Fe(CN)6, 0.5 mM K4Fe(CN)6, 0.1% Titon X-100, 0.1% DMSO. After 12 h incubation at 37°C intact seedling were photographed. Alternatively, stained seedlings were vacuum infiltrated with Tissue-Tek OCT compound prior to freezing in liquid nitrogen. A bright cryostat microtome (model 5030) was used to cut 20 µm sections at -23°C.

Other modifications of the present invention will be apparent to those skilled in the art without departing from the scope of the invention.

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#### **Claims**

1. A promoter comprising the DNA sequence of an oil seed rape cysteine protease gene promoter of class 1, 2 or 6.

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- 2. A promoter comprising at least part of the DNA sequence as shown in Figure No.19, or at least part of a sequence that has substantial homology therewith, or a variant thereof.
- A promoter comprising at least part of the DNA sequence as shown in Figure No. 20 or at least part of a sequence that has substantial homology therewith, or a variant thereof.
- 4. A promoter comprising at least part of the DNA sequence as shown in Figure 21 or at least part of a sequence that has substantial homology therewith, or a variant thereof.
  - 5. A promoter having the characteristic motif or features of a promoter of any one of the preceding claims.
- A recombinant DNA construct comprising the promoter of any preceding claim operably linked to a gene of interest.
  - 7. A recombinant DNA construct functional in a plant comprising a disrupter gene encoding a product capable of disrupting cell function, and a promoter according to any preceding claim for the disrupter gene, the disrupter gene being functionally linked to and controlled by an externally regulatable gene control region which includes a promoter which is inducible by the external application of a chemical inducer.
- 8. A recombinant DNA construct according to claim 7 wherein the inducible promoter is

  functionally linked to and controls a repressor protein gene and in which the disrupter
  gene promoter includes an operator sequence which is recognised by the repressor

protein, so that in the presence of the inducer the repressor protein is produced which interacts with the operator sequence thereby disabling the second promoter and inhibiting expression of the disrupter gene.

A recombinant DNA construct according to claim 7 or claim 8 wherein the disrupter gene is a nucleotide sequence, which is in sense orientation to an endogenous plant gene which is essential to plant development or a gene conferring a desired characteristic on the plant, or comprises a partial sense sequence of the endogenous plant gene.

10. A recombinant DNA construct according to claim 7 or claim 8 wherein the disrupter gene is a nucleotide sequence, which is in antisense orientation to an endogenous plant gene which is essential to plant development or a gene conferring a desired characteristic on the plant.

- 11. A recombinant DNA construct according to claim 9 or claim 10 wherein the endogenous plant gene is essential to seed germination or early seedling development.
- 12. A recombinant DNA construct according to any one of claims 7 to 11 wherein the
  externally regulatable gene control region is a chemically inducible gene promoter
  sequence from the glutathione S-transferase system, the Alc system or the ecdysone
  system.
- 13. A recombinant DNA construct according to any one of claims 8 to 13 wherein the repressor protein gene encodes a bacterial repressor.
  - 14. A recombinant DNA construct according to claim 13 wherein the repressor protein gene encodes the *lac* repressor or a repressor used by 434, P22 or lambda-bacteriophages.

10

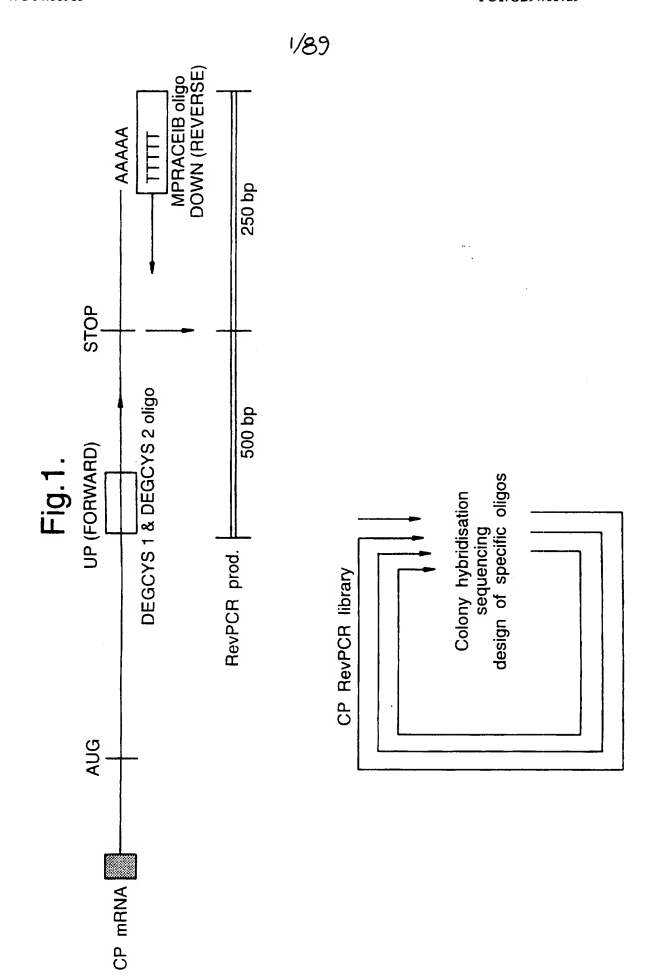
- 15. A recombinant DNA construct according to any one of claims 7 to 14 wherein the disrupter gene or disrupter promoter contains a "pseudo-operator".
- 16. A recombinant DNA construct according to any one of claims 7 to 15 wherein the disrupter gene is a cytotoxic gene.
  - 17. A recombinant DNA construct according to any one of claims 7 to 15 wherein the disrupter gene encodes a recombinase or transposase adapted to excise a nucleotide sequence flanked by recombinase recognition sequences.

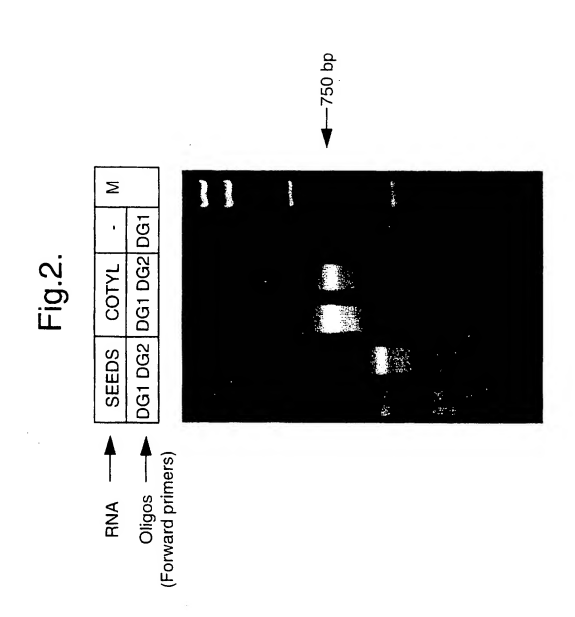
5

- 18. DNA comprising at least part of the sequence shown in Figure 12 or at least part of a sequence that has substantial homology therewith or a variant thereof, and which codes for a cysteine protease.
- DNA comprising at least part of the sequence shown in Figure 13 or at least part of a sequence that has substantial homology therewith or a variant thereof, and which codes for a cysteine protease.
- DNA comprising at least part of the sequence shown in Figure 14 or at least part of a sequence that has substantial homology therewith or a variant thereof and which codes for a cysteine protease.
  - 21. DNA comprising at least part of the sequence shown in Figure 15 or at least part of a sequence that has substantial homology therewith or a variant thereof, and which codes for a cysteine protease.
  - 22. DNA comprising at least part of the sequence shown in Figure 16 or at least part of a sequence that has substantial homology therewith or a variant thereof, and which codes for a cysteine protease.

- 23. DNA comprising at least part of the sequence shown in Figure 17 or at least part of a sequence that has substantial homology therewith or a variant thereof, and which codes for a cysteine protease.
- A recombinant DNA construct functional in a plant comprising the DNA of any one of claims 18 to 23 operably linked to a promoter.
- 25. A recombinant DNA construct according to any one of claims 6 to 17 and 24 wherein the construct is capable of being expressed in the tissue or tissues of a germinating seedling or a developing grain or a plant when the construct is integrated, preferably stably integrated, within the grain's or seedling's or plant's genomic DNA.
  - An expression system for the tissue or tissues of a plant material, the expression system comprising a gene of interest fused to a gene promoter as defined in any one of claims 1 to 5 wherein the expression system is capable of being expressed in the tissue or tissues of the plant material.
- An expression system according to claim 26 wherein the expression system is for at least the tissue of a germinating seedling or developing grain or plant (eg in the root, cotyledons, leaves and stem).
  - 28. An expression system according to claim 26 or claim 27 wherein the expression system is integrated, preferably stably integrated, within a germinating seedling's genomic DNA or a developing grain's genomic DNA or a plant's genomic DNA.
    - 29. An expression system comprising a construct according to any one of claims 6 to 17 and 24.

- A recombinant plant genome comprising a promoter of any one of claims 1 to 5, DNA of any one of claims 18 to 23, a recombinant DNA construct of any one of claims 6 to 17 or 24 or an expression system according to any one of claims 25 to 29.
- 5 31. A plant, plant seed or plant cell having a recombinant plant genome of claim 30.
  - 32. Protected plant germplasm comprising a plant comprising a recombinant DNA construct of any one of claims 6 to 17 or 24.
- 10 33. A plant or seed which is incapable of growing to maturity comprising a recombinant DNA construct of any one of claims 6 to 17 or 24.
  - 34. The use of a gene promoter as defined in any one of claims 1 to 5 to induce expression of a gene of interest when fused to the gene promoter in the tissue or tissues of a plant material.
- 35. The use according to claim 34 wherein the gene promoter is used to induce expression of a gene of interest when fused to the gene promoter in at least the tissue or tissues of a germinating seedling or a developing grain or a plant (eg in the root, cotyledons, leaves and stem).
  - 36. A promoter, a construct or an expression system substantially as hereinbefore described with reference to any one of Figures 12 to 17 and 19 to 21.





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sednences:
Fig.3A. 1/8
                                                                                                                                                                                                                        (1-501)
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                                                                                                                                                                                                    1-468
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                              Identity
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                                                                                                                                                                                57% humd
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                                                                                                                                 H
                                                                                                                    Matching-weight
NUCLEIC-Res-length
                                       Output line length
                            Nucleic Alphabet
                                                                                         AMINO-Res-length
         Solution Parameters:
                                                                                                  DELetion-weight
                                                                                                                                                                               OSR8.401COD
                                                                                                                                                                                                                                 OSR8.389COD
                                                                                                                                                                                                                                            OSR8.387COD
                                                                                                                                                                                         OSR8.406COD
                                                                                                                                                                                                   OSR8.403COD
                                                                                                                                                                                                              OSR8.404COD
                                                                                                                                                                                                                        OSR8.402COD
                                                                                                                                                            Clustered order of
                                                                                                            LEngth-factor
                                                                                                                                         SPread-factor
                                                                     Randomization
                                                           Histogram
                                                  Compress
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15.
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127/474 13.
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Fig.3A. 2/8 CYS 8.407MR

Region Alignment: (listed in OSR8.401CO)       1       NNNNNCaatNg         OSR8.406CO       6       1       NNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNN
Region Alicoosr8.401C0 OSR8.406C0 OSR8.403C0 OSR8.402C0 OSR8.389C0

5/89

		<b>TIG.3A.</b> 3/8 CYS 8.401
OSR8.401CO	62	ACACaGAGAAGATTATCCTTAtCaaGaacgtgATGGcac
OSR8.406CO	62	Acaccadoladalpadicirracqqeqetecaaried aaaarecaaftrorracti
OSR8.403CO	62	52 ACACCGAGAAAGACTACCACGGAACCAATGG CAAATGCACTTTAACTTAA CHILLIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIII
OSR8.404CO	62	ACACCGAGAAAGACTATCCTTACCACGGAACCAATGG CAAATGCAACTTTACTTAA (
OSR8.402CO	62	ACACCGAGCAAGATTAATTACGGTGGTTACCAATGG MAATGCAATTATTACTGAA
OSR8.389CO	62	ACACCGAGCAAGATTATCCTTACGGTGGTTCCAATGG AAAATGCAATTCTTTACTGAA
OSR8.387CO	62	ACACCGAGCAAGTTALTACGGTGGTTCAATGG BAAATGCAATTACTTTACTGAA
consensus		ACACcGAGcAAGAtTAtCCTTAcCgtGgttccaATGGcacaaaatgcAAttcTttacTgaa

Fig.3A. 4/8	OSR8.401CO 120 GAATagAAagGTTGTgACaATTGATaGcTACGctGGTGTaaaatCaaAtGAcGAgAaaGCC	OSR8.406CO 117 GAATTCAAGAGTTGTAAGTTATOGATGGETACGAAGATGTTCCTACTGAAGATGAAGGGCC	OSR8.403CO 120 GAATTCAAGAGTTGTAACCATGCATACCAAGATGTTCTAGTAAGATGAAACCGCG	OSR8.404CO 120 AAATTCGAGACTGTGTGTGTGTGTGTGTGTGTGTGTGTGT	OSR8.402CO 120 GAATTCAAGAGTTGTAACTATTGATGGATACGAAGATGTTCCTAGTAAGATGAACCDCD	OSR8.389CO 120 GAATTCAAGAGTTGTAATGATGGTTACGAAGATGTTCCTACTGAAGATGATGAAGATGTTCCTACTGAAGATGAAGATGAAACGGCG	OSR8.387CO 120 GAATTGAAGAGTTGTAATGGATGGGTACGAAGATGTTCCTACTGAAGATGATGAAGATGAAAAAAAA	consensus gAATtcaAgaGTTGTaACtATtGATgG-TACGaaGaTGTtcctactaAaGAtGAaaccqcc
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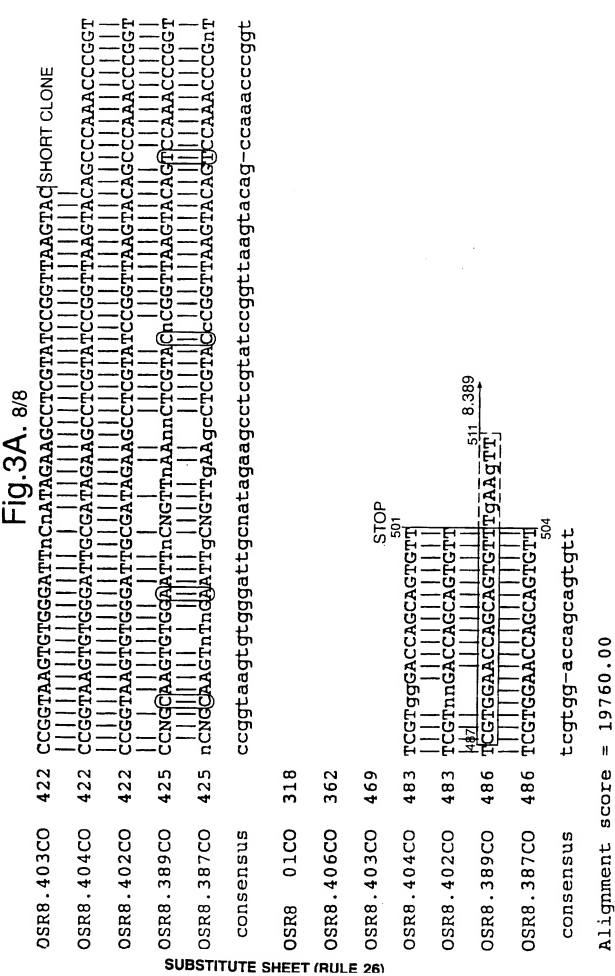
Fig.3A. 5/8

		O 181 TIGAGAGAGAGAGTITCATACCAGCCTGTGAGTGTTGCTATTGATGCTGGAAGAGCTTT					s ttgaagagagcagtttcataccagcctgtgagtgttgctattgatgctggtggaagagctT
OSR8.401CO	OSR8.406CO	OSR8.403CO	OSR8 404CO	OSR8.402CO	OSR8.389CO	OSR8.387CO	consensus

Fig. 3A. 6/8 CYS 8.407R 3'	CTGGcccaTGTtc	239 TCCAACATTACCAATCGGGATCTTCACTGGGAAGTGTGGGAAGAATGTGGAATCAAGAGAT	TTCACTGGAAAGTGTGGTACGAATATGGATCZ	ACCAATCTGGAATCTTCACTGGAAAGTGTGTACGACTATGGATCACG	242 TCCAACATTACCAATCTTCACTGGAAAGTGTGGTACGAATATGGATCACGCTGT	CATTACC	242 Incaachtraccaathgegarctrcacteggaagtereggaagtereggaagtereggaagtereggaagte	TcCAacatTACcaatcgGGaATcTTCaCTGG-aagTGTgg-ACaaat-TgGAtCAcGCaGT	CGAGTAGCATCC 303 GCTCATCGTAGGATA SHORT CLONE	GGCOCTTGCTTATGCOTCAGAGGAGCGCOTTGACTTGGATTGTAGCGAACTC		TGGTTATGGATCAGAGGAGCGtGTTGACTATTGGATTGTACGTAACTC	
	OSR8.401CO	OSR8.406CO	OSR8.403CO	OSR8.404CO	OSR8 402CO	OSR8.389CO	OSR8.387CO	consensus	OSR8.401CO	OSR8.406CO	OSR8.403CO	OSR8.404CO	OSR8.402CO
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8// . <b>C</b> .5: -	GGTGGGTGTTGGTTATGGTTCAGAGGACGGTATTGACTATTGGATTGTAA	I	TAtggttcagagaacggt-ttgactattggattgta-g-aactctt			ATTAGGATGGAGAGAAACGTG			ATTAG	ATTAGAATGGAGAGAAAQ	gagaggatggttacattaggatggagagaaacgtg		
			ggTggc-GTtGGt1		<b>U</b> -	GGTACACGTTGG	GGTACACGTTGG	GGTACACtTTGG	GGTACACGTTGG	GGTACACGTTGG	ggtacacgttgggg		
	303	303		318	357	364	364	364	364	364		318	357
	OSR8.389CO	)SR8.387CO	consensus	OSR8.401CO	OSR8.406CO	)SR8.403CO	)SR8.404CO	OSR8.402CO	OSR8.389CO	)SR8.387CO	consensus	OSR8.401CO	OSR8.406CO
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CTAAGAUT

CTTGGGAGITTTATA

ATAATCA

8.387

Criggagnnitara

ATAATCA

Fig.3B. 1/3 1.00 5.00 11 11 11 Ħ NUCLEIC-Res-length Matching-weight DELetion-weight LEngth-factor SPread-factor

Clustered order of selected sequences:

14. OSR8.389NCOD 75% humd. (1-240) 17. OSR8.387NCOD with 8.402 (1-246) 7. OSR8.402NCOD 8.404 (1-242) 6. OSR8.404NCOD 8.403 (1-242) (listed in Clustered order Alignment: Region

TGAAGTTGTTTTAAAATAAAAACTCA TGAAGTTHTTTTAAAATAAAACTCA OSR8.389NC **OSR8.387NC** 

OSR8.387NC OSR8.402NC OSR8.404NC

consensus

GTAACAAAAAGAATCTCATGCAGTAATCAAATTGGGATTGTTATAA| GTTAAAT GTAACAAAAAG@ATCTCATGCAGTAATCAAATTGGGATTGTTATAA GTTAAAT TGAAG TGAAG STOP

TGAAGttgT---AAAA--aA-CTCAtgca-TAATCAa-TTGGGA-T-TtATAAc---A-

Fig.3B. 2/3

Fig.3B. 3/3	9NC 176 THANTITES FIGHT ATTENTA SAGA SERVITA A PARA PARA PARA PARA PARA PARA PARA	
	OSR8.389N	

				13/8	)				
176 TTAATTTGG TTGTTTATGTATTAAGAGGTATAAT AAAATGATATATTTCTGTTAA	TTGTTTATGTATTAAGAGTAGTAAT		TTGCTTTG@CTTGATTATGTATTAAGAGAAATA	poly A signal TTTTTGgcTTG-TTATGTATTAAGAGAA-TATAATaAAAATGaTATATTCTCttAaca	AAAAAA	tcar	1 GCAAAAAAA 	1 GCAAAAAAA TTTTTTTTT	gcaaaaaa
17	176	172	173		233	235	231	231	
OSRB.389NC	OSR8.387NC	OSR8.402NC	OSR8.404NC	consensus	OSR8.389NC	OSR8.387NC	OSRB.402NC	OSR8.404NC	consensus
	30	09111	UIE	SHEET	HULE	26)			

Fig. 4. 1/4	
	: (CLASS 1)
	317:
	1 to
	from
	.401COD
	OSR8.

nnnnncaatn ggggnntgat ggacnnnnnt tttcaatttg tcattaaaaa ccatgggatt gacacagaga 9 50 30 20 10

ttqtqacaat ttatcaagaa cgtgatggca cctgtaagaa agataagttg aatagaaagg 130 120 110 100 aaqattatcc

ttactagaag ctgtagncgc 200 190 gctggtgtaa aatcaaatga cgagaaagcg 180 170 160

220 230 240 250 250 260 270 280 agtgttggtag cgagagagag tttcagttat actctaaggg aatattctct ggcccatgtt

14/89

tcagccagtt

290 300 300 caacatcatt ggaccacgca gtgctcatcg taggata

GEL: pri 40

Range to print (<CR>=ALL): OSR8-402 from 1 to 743: (CLASS2)

aacaccgagc tcatgaaaaa cggcggtttg 09 50 tttcaattca gggtgcaacg ggngactgat ggactatgct 30

tgatagctac

Fig. 4. 2/4

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	140 ttgtaactat	210 ccagcctgtg	280 ggaaagtgtg	350 attggattgt	420 ggcgtctaaa	490 gttcgtnnga	560 aagttaaatt	630 ttaatctgta	700 gtattaagag	
	130 aattcaagag	200 cagtttcata	270 aatcttcact	340 ggcgttgact	410 agagaaacgt	480 cccaaacccg	550 ggattgttat	620 ccatagggat	690 gcttgattat	·
<b>t</b>	120 tttactgaag	190 ttgaagagag	260 accaatctgg	330 gtcagagaac	400 attaggatgg	470 ttaagtacag	540 aatcaaattg	610 aattgattca	680 atttgctttg	<b>ਕੈਕ</b> ਕੋ
17	110 aatgcaattc	180 tgaaaccncn	250 ttccaacatt	320 ttggttatgg	390 ggatggttac	460 tcgtatccgg	530 ctcatgcagt	600 tttcgaaaaa	670 ttcaaagaat	740 agcaaaaaa
	100 tccaatggaa	170 ctagtaaaga	240 tggaannnct	310 gtggtggcgg	380 tttggggaga	450 gatagaagcc	520 caaaaagaat	590 atgtatggta	660 tatcatttca	730 atttctcagc
	90 ttaccgtggt	160 gaagatgttc	230 ttgatnctgg	300 ggatcacgct	370 tggggtacac	440 gtgggattgc	510 ttgaaggtaa	580 tattgtttgt	650 tgttggtcaa	720 aaaatgatat
	80 aagattatcc	150 tgatggatac	220 agtgttncta	290 gtacgaatat	360 acgtaactct	430 tccggtaagt	500 ccagcagtgt	570 aatcttgtat	640 taaatctcta	710 aaatataata

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to 744: (CLASS 6)

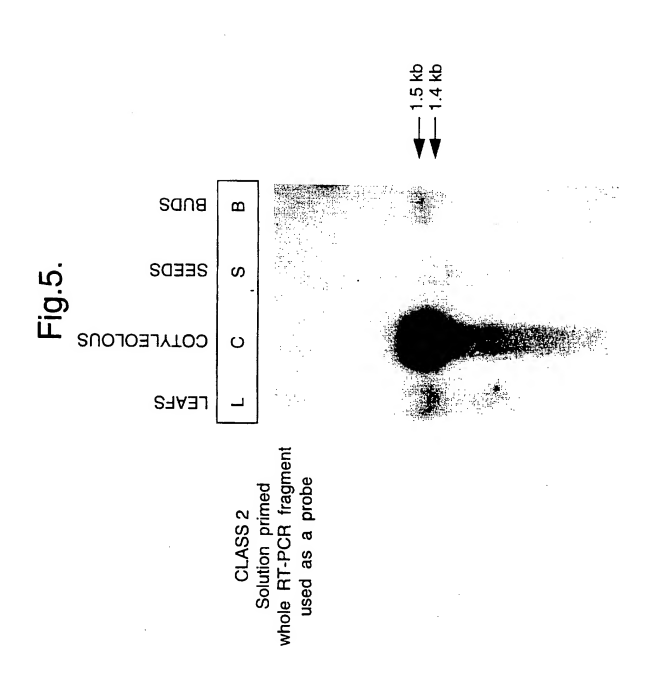
from 1

CYS8-389

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60	130	200	270	340
cggcggtttg	aattcaagag	cagtttcata	gatcttcact	ggtattgact
50	120	190	260	330
tcatgaaaaa	tttactgaag	ttgaagagag	accaatcggg	ttcagagaac
40	110	180	250	320
tttcaattca	aatgcaattc	tgaaacggcg	ttccaacatt	ttggttatgg
30	100	170	240	310
ggactatgct	tccaatggaa	ctactgaaga	tggaagagtt	gtggtggctg
20	90	160	230	300
gggggttgat	ttaccgtggt	gaagatgttc	ttgaagctgg	agatcatgca
10	80	150	220	290
gggtgcaacg	aagattatcc	tgatggttac	agtgttgcca	ggacaaatct

16/89

	420 ggcaaggtcc	490 ccggttcgtg	560 aactaagatt	630 tacggattta	700 gtttatgtat	
	410 agagaaactt	480 cagtccaaac	550 ggagttttat	620 tgatccacca	690 taatttggtt	
	400 attagaatgg	470 cggttaagta	540 taatcacttg	610 agaaggtatt	680 aaagaaagat	aaaa
Fig.4. 4/4	390 ggatggttac	460 nnctcgtacn	530 taaaactcaa	600 atatcaaaaa	670 catttcgttt	740 tcttaaaaaa
	380 gttggggaga	450 tncngttnaa	520 ttttttaaaa	590 tatgtatagt	660 cgatcaatat	730 gatatatttc
	370 tggggtacac	440 agtgtggaat	510 tgtttgaagt	580 ttattgtttg	650 atccttatgt	720 ataataaaat
	360 aaggaactcg	430 aagtccngca	500 gaaccagcag	570 taatctcata	640 atctgtatgg	710 taagagaagt



19/89

**ORGANS** ≥ LIGHT Ω 9 Ŋ DAYS AFTER IMBIBITION 9  $\mathbf{O}$ 4 Fig.6. DARK က  $\sim$ 0.5 0 MS CTRL MS CTRL CODING NEG. CLASS 1 CDCYS12 PCR PROBE CLASS 6 CDCY66 CLASS 2 CDCY25 NON CODING NEG. PCR PROBES

Fig.7. 1/4

			†/- · · · · · · · · · · · · · · · · · · ·
	CYS4_BRANA	123 sc	123 scwafstaaaveginkivtgelvslaegelvdcdksynqpcngglmdyaFQFIMKNGGI
0.00010	OSR8.403CO	H	XXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXX
	OSR8.404CO	н	
_	OSR8.402CO	н	GCNGXLMDYAFQFIMKNGGI
	OSR8.389CO	1020/ 1000	GCNGGIMDYAFQFIMKNGGI
CLASS 6	CLASS 6 OSR8.387CO	30 % Hulliu. 1	GCNGGLMDYAFQFIMKNGGI
_	OSR8.406CO	1 20%	WITH 8.103 XXXXXXXXXXXXXXFQFIMKNGGI
CLASS 1	OSR8.401CO	jez% numa. 1	XXnXgXmdXXFQFVJKNhG①

scwafstaaaveginkivtgelvslseqelvdcdksynq--n--lmdy-FQFimKNgGI

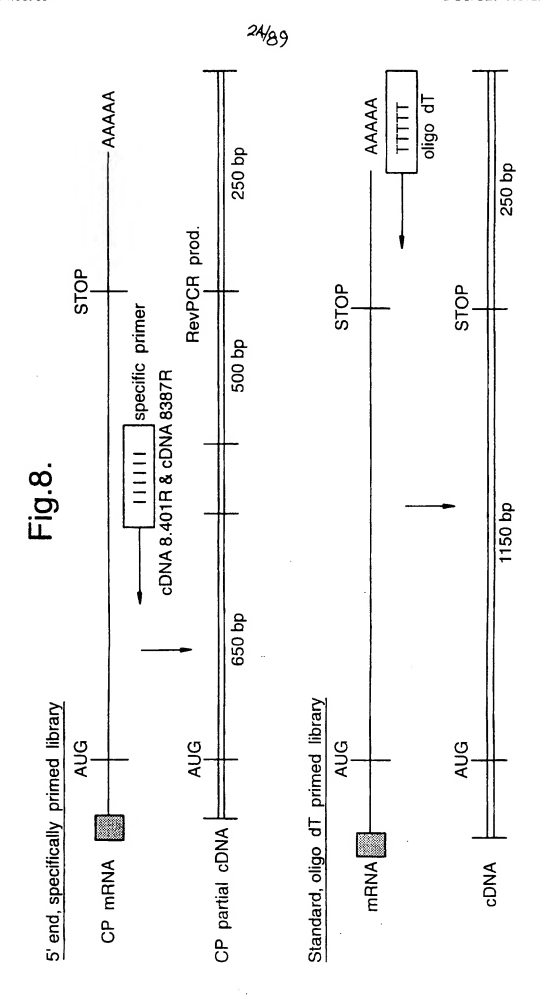
consensus

-ig.7. 2/4

Fig. 7. 3/4

CYS4_BRANA OSR8.403CO	244	HYOSGIFTGKCGTNMDHAVVAVGYGSENGVDYWIVRNSWGTRWGEDGYIRMERN VASR 
OSR8.404CO	83	HYOSGIFTGKCGT(HMDHAVVAVGYGSENGVDYWIVRNSWGTRWGEDGYIRMERN VXSK
OSR8.402CO	83	HYQSGIFTGKCGTNMDHAVVAVGYGSENGVDYWIVRNSWGT()WGEDGYIRMERN VASK
OSR8.389CO	83	HYOSGIFIGECGINCDHAVVAVGIGSENGIDYWIVRNSWGTRWGEDGYIRMERNCARSER
OSR8.387CO	83	KYOXGIFTGKCGTNLDHAVVAVGYGSENGIDYWIVRNSWGTRWGEDGYIRMERNLARBK
OSR8.406CO	82	1
OSR8.401CO	83 (	TYERGIFGCPONE
consensus	•	hYqsGIFtGkCgTn-DHAVvaVGygsengvdywivrnswgtrwgedgyirmernlvask

FIG. / . 4/4 328	KCGIaIEASYPVKYspnpvrgtssv	KCGIXIEASYPVKYSHORT CLONE	KCGIAIEASYPVKYSPNPVRGTSSV		CONTROL   CONT				kcgiaieasypvkyspnpvrgtssv 168	= 3484.00 STOP
	304	143	143	143	144	144	121	106		score =
	CYS4_BRANA	OSR8.403CO	OSR8.404CO	OSR8.402CO	OSR8.389CO	OSR8.387CO	OSR8.406CO	OSR8.401CO	consensus	Alignment :



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2
1.00
                        5.00
                                 17
        11
                 11
                        11
                                NUCLEIC-Res-length
AMINO-Res-length
       DELetion-weight
                        Matching-weight
                LEngth-factor
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Fig.9. 1/2

sednences: selected Clustered order of SPread-factor

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(1-328)
(1-225)
(1-230)
(1-225)
\begin{array}{c} \text{CYS4} & \text{BRANA} = \text{COT 44} \\ \text{CYS2} \overline{\text{UP6}} & 1 \\ \text{CYS2} \overline{\text{UP7}} & 1 \\ \text{CYS2} \overline{\text{UP8}} & 2 \\ \text{CYS2} \overline{\text{UP8}} & 2 \\ \end{array}
```

Region Alignment: (listed in Clustered order)

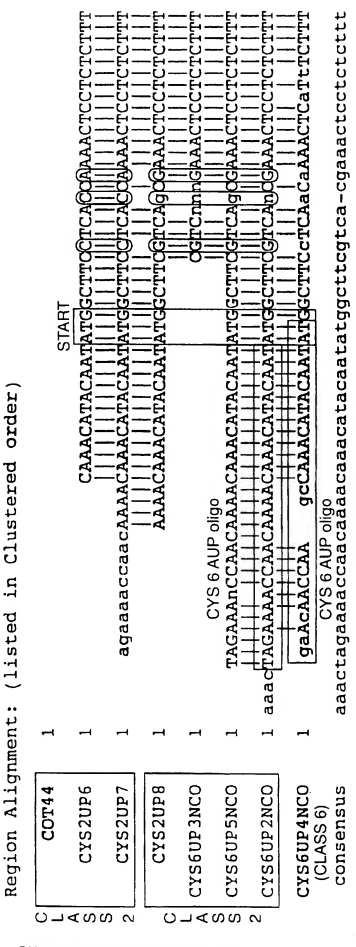
ATG		- FROTROMASSPRILLING FILL FOR STANDING FILL FILL FILL FOR STAND FO		$\tt rkptkqtynmasspkllslllyvfvslasgyesiisdnhlslpsdrswrtdeev-SIYLr$
-	-	₽.	-	
CYS4_BRANA	CYS2UP6_1	CYS2UP7_1	CYS2UP8_2	consensus

Fig.9. 2/2

IGKSNSNSN 	WSLEHGKSNSNSN	2 WSLEHGKSNSNSINOODERFNIFKDNLRFIDLHNENNKNATYKLGLTIFADLTNDEYR	WSLEHGKSNSNSN	WSLEHGKSNSNSNGIINQQDERFNIFKDNLRFIDLHNeNNKNATYKLGLTIFAdLTNDEYR	SLYLGARTEPVRritkaknvnmkysaavnvdevpytvdwr@kgavnaikDQGTCC	SLYLGARTEPVRXXTKAXNVNMKYSAAVNXVEVPETVDWRKKGAVNAIKDQGTCGSCWAFS	3 SLYLGARTEPVRXXTKAXNVNMKYSAAVNXVEVPETVDWRKKGAVNAIKDQGTCGSCWAFS	SLYLGARTEPVRXXTKAXNVNMKYSAAVNXVEVPETVDWRGKGAVNAIK@QGGCC	SLYLGARTEPVRxxTKAxNVNMKYSAAVNxvEVPeTVDWR-KGAVNAIKdQGtCGSCWAFS	TAAAVEGINKIVTGELVSLSEQELV		TAAAVEGINKIVTGELVSLSEQELV	TAAAVEGINKIVTGELÜSLSEQELV	TAAAVEGINKİVTGELvSLSEQELVDCDKSYNQGCNGGIMDYAFQFimkngglntekdypy
	57	62	58		9	118	12	119		129	179	184	180	
CYS4_BRANA	CYS2UP6_1	CYS2UP7_1	CYS2UP8_2	consensus	CYS4_BRANA	CYS2UP6_1	CYS2UP7_1	CYS2UP8_2	consensus	CYS4 BRANA	CYSZUP6_1	CYS2UP7_1	CYS2UP8_2	consensus

Fig.10. 1/6	= 2 = 1.00 = 0 = 5.00 = 90	selected sequences:	(1-1102) (1-675) (1-691) (1-679) (1-322) (1-307) (1-288) (1-255)
Sequencing results	AMINO-Res-length DELetion-weight LEngth-factor Matching-weight NUCLEIC-Res-length SPread-factor	stered order of sel	9. COT44 10. CYS2UP6 11. CYS2UP7 12. CYS2UP8 3. CYS6UP3NCOD 1. CYS6UP5NCOD 4. CYS6UP2NCOD 2. CYS6UP4NCOD

Fig. 10. 2/6



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CYS2UP7	118	CATCTCAGICTTCCATCTGACCGITCGAGAGAACCGATGAAGAAGTGAIAIIIIIIIIII
CYS2UP8	107	107 CATCTCATITCTTCCATCTGACCGTCGAGAACCGATGAAGAAGTGAAGTGATGTCTACT
CYS6UP3NCO	84	CATCTCATECTTCCATCTGACCGCTCGAGAGCCGATGAAGAAGTGATGTCCATCTACT
CYS6UP5NCO	119	CATCTCATECTTCCATCTGACCGCTCGTGGAGAACCGATGAAGAAGTGATGTCCATCTACT
CYS6UP2NCO	123	CATCTCAUTCTTCCATCTGACCGCTCGTGGAGAACCGATGAAGAGAAGTGATGTCTACT
CYS6UP4NCO	115	CATCTCAATCTTCCATCGGACnGCTCATGGAGAACCGATGAAGAAGTGAGGTCCATCTACT
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COT44	CYS2UP6	CYS2UP7	CYS2UP8	CYS6UP3NCO	CYS6UP5NCO	CYS6UP2NCO	CYS6UP4NCO	consensu

Fig.10. 6/6

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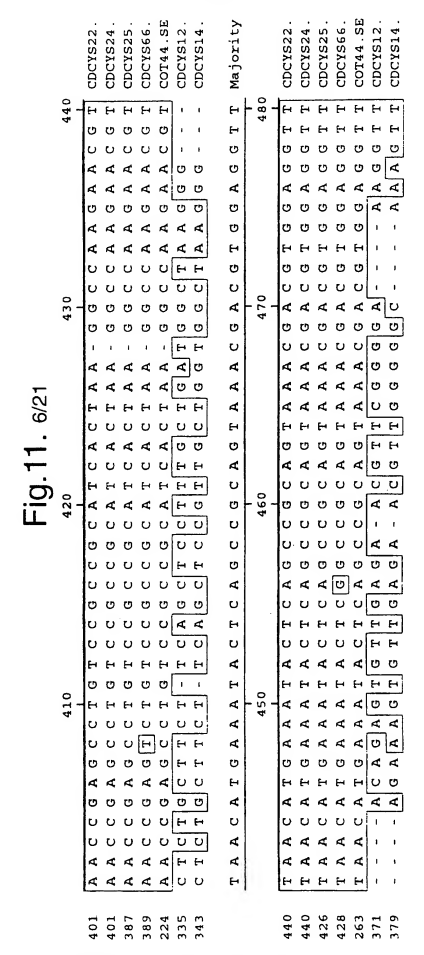
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Fig. 11. 7/21

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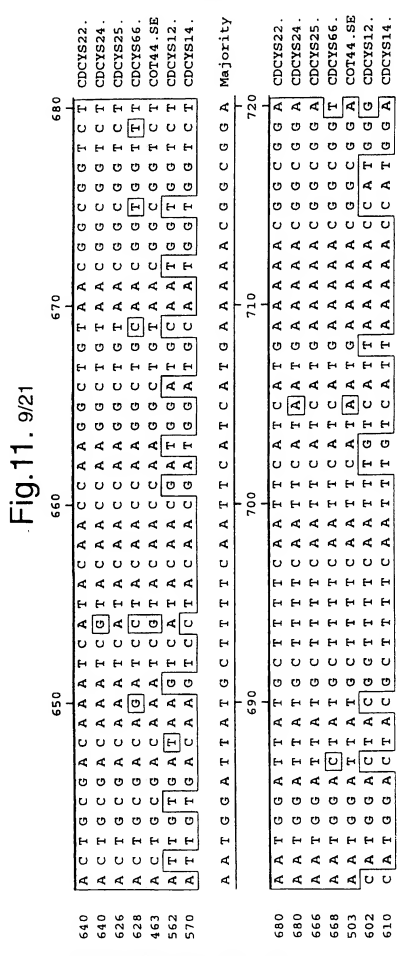


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Fig. 11. 12/21	890 1	ATTGATGCTGGTGG	ATTGATGCTGGTGGAAGAG	ATTGATGCTGGAAGCTTTCCAACATACCAA	ATTGATGGTGGAAGGGTTTT	ATCTGTGGGAGAGAGCGTTTCAGTTACTCT	ATCTGTGGC	G G A A T C T T C A C T G G A A G T G T G G T A C G A C T A T G G A T C Majority	930 940 950	GAATCTTCA	GAATCTTCACTGGAAAGTG	GAATCTTCACTGGAAAGTGTGGTACGA	GENT CIICACIGGAAGTGTGGACAAA	CANICILCACTGGAAAGTGTACGAAAT	G A A I A I I C I G G C C C A I G I I C A A	GAATATTCI
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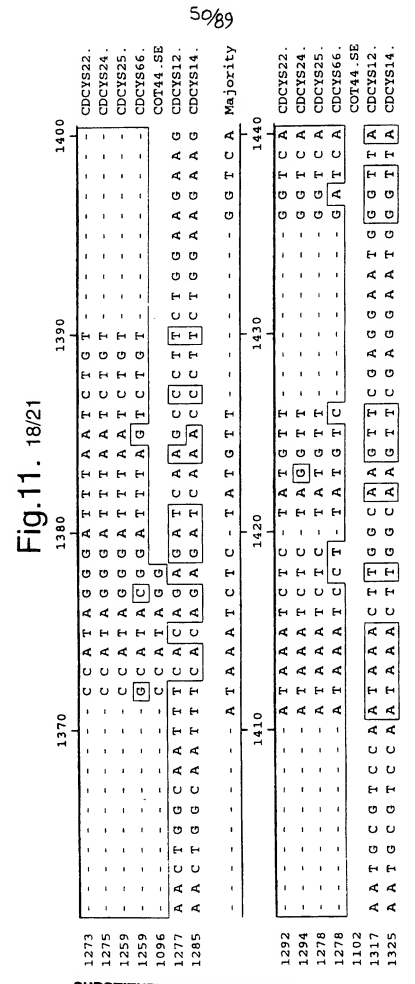
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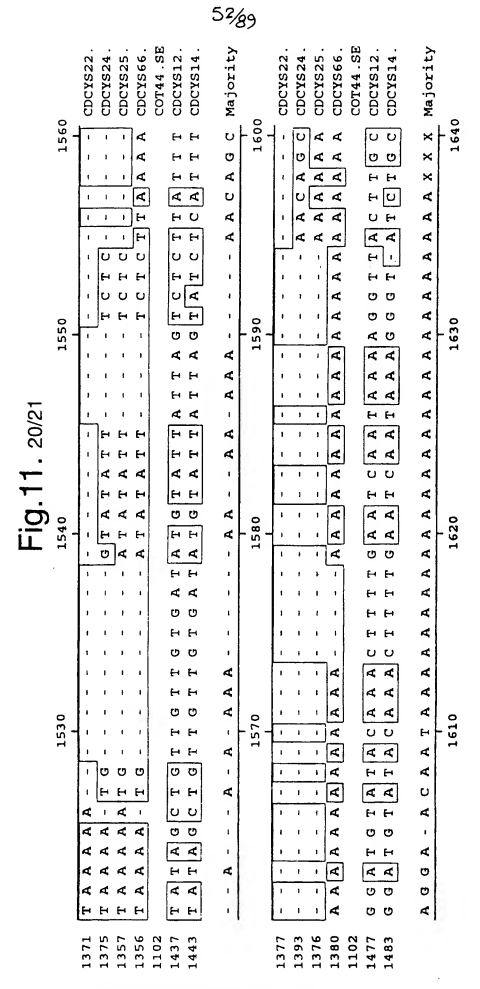
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Decoration 'Decoration #1': Box residues that match the Consensus exactly.

	9		6111 60	TCCG 120	TCGG 180	CGAC 240	ACTC 300	CTGA 360	TGGA 420	TCTT 480	:AGCC 540	.GGTC 600	16ATT 660	GTGA 720	GTAG 780
Fig. 12. 1/2		1	TCCTTTGTTT	TCATCTTCCG	TACGCTTCGG	GICACACGAC	GATCTGACTC	CCTTTGCTGA	GTTGATTGGA	TGTTGGTCTT	CTCATCAGCC	AATGGTGGTC	GAGAAAGATT	AAGGTTGTGA	GAAGCTGTAG
	90	Tirrition	ATCGTCTTCT	CTTCTCATCA	CGGCAAAACG	TCACGACTTC	TGCCTTCGCG	TTCTTCAGCT	TCCAGATICT	CTGCGGAGCG	AACAGGAGAT	CGATGGATGC	GATTGACACA	GTTGAATAGA	AGCGTTACTA
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	. 20	Terretorn	TTTCCCTGTA	CTICICCCTT	CGAGCTGTTC	ACAACACAGG	CGCTAACTCT	CAAGGCCTCT	ACAGAGTGTT	AGCTGTTACT	TGGAGCTATG	AGAACTCATT	CGCTTTTCAA	AGAACGTGAT	CTACGCTGGT
	10	Littin	CITGITITEG	CTCTCACTTT	ATGACATCTC	AGGAAGAGAG	ACAACAACAT	ACCACGAGII	TGGCTAAGGG	GGAAGAAAGG	TCTCGGCGAC	TCTCTGAGCA	TCATGGACTA	ATCCTTATCA	CAATTGATAG
		5	SUBS	STIT	UTE	SH	EET	(RL	JLE	261					

# Fig. 12. 2/2

500 1260 TTATACTCTA AAGAAGAATG TACTITATA TTTGAATCA AAAAAAAAA GTATGCGGAA GTTGCTGTG ACCACCAGAA AAGAGITGGG CCGICCCCIC rctgctgtgt ATCGTAGGAT I TCAAAC TC AGCGTTTCAG AAAAAAAAA CGCAGAAGGA GCCCTTCTGG CTCTTGGGGA CCCTCCTCCA CGAGCTAGAG **IGAGAGATGA** GTATACAAAC CGCAGTGCTC CGTCTGTGAT **I G A T G A G A C T** AGAGGAAGTI GGAGCGAGAG AAAAAAAAA CAGAGATCAA CATTGGACCA TCGTGAAGAA ACACCGGCAA CACATCCAAA GGAAATGCTG GGGTTATGTA ATCTATAAGC TTATTTGGAT GTGATTACCC ATTGTTCAGC GGTATCTGTG ACTIGCAGGA CACAATAAAA GGCAATTTCA TATTAGTCTC ATGCAGCGTA GATTACTGGA CCCATCAAGA TGTTTCTCGT TGTTGTCCGC TCGAGGAAT TGTTCAACAT GGATTCGTTT CITITCACCI AAAAAA 1577 AGTTAGTGTT GTTTGGTTTG ACTTGGCAAG TGATATGTAT CTCTGGCCCA GITTAICCAC GGCTICATAT TAAGAAACT GAACGGTGTT GAAATGCAAC TGGTCGTCAT AAGCCTCTTT AAAAAAAAA CGGCTCAGCC ACGGTTCAAA GAATGGATGG CGTCCAATAA GCTGTTGTTG GTTGCAAGGA ATAAAAGGTT AGGGAATATT **CCAACATGCT** CCGGCCCCAC CGAGAAACTI GTCTTTGCCT T T C A C A C G G T

Fig.13. 1/2

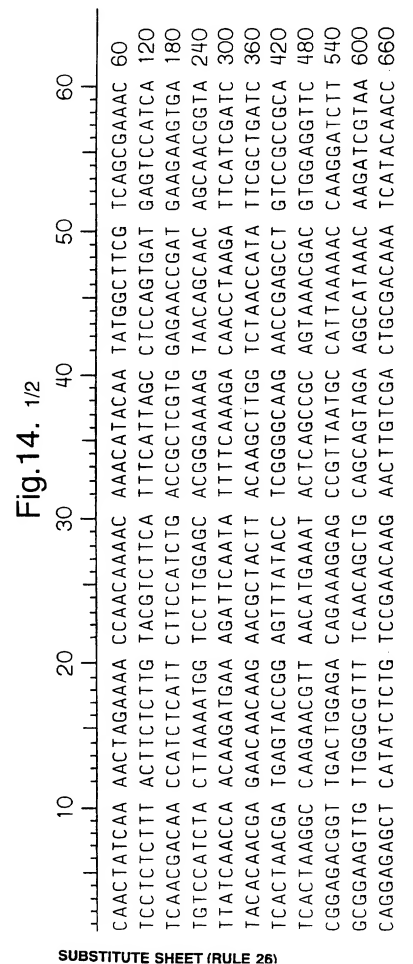
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			9	120	180	240	300	360	420	480	240	009	099	720	780
	09	1111111111	CATCGICITC	GCTTCCCATC	ACGGCAAAAC	ATCACGACTT	ATGCCTTCGC	CTTCAGCTCC	CAGATICIGI	GCGGAGCGTG	CAGGAGATCT	ATGGATGCAA	TCGACACAGA	TGAAAAGAAA	CGTTACTGGA
	90	ATGTCACCAT	TCTTCTCA	TGCCAGAGAC	TTTAGAGACA	CTCTCACTCA	CTCTCTGCTT	GGCAAAGTTC	CAAGGAAGCT	CAGATTGTAA	TCCTACAACG	AACCATGGAA	AAAGATAAGT	GACGAGAAAG	
	04	Listerin	AGGAAAAGAA	TCTTCTAGTT	CGACGCTTGG	GATTGAAATC	TACTTACTCT	TCGTCTTGGA	GAACGTTGGG	TGTCAAAGAT	AGGAATCAAC	TTGTGACAAG	TGTCATTAAA	CACCTGTAAA	AAAATCAAAC
	30		TTTCCCTGTA	TCTTCTCCCT	CCGAGCTGTT	GACAACACAG	TCGCTAACTC	TCAAGGCCTC	AAAGTGTTGA	CTGTTACTAA	GAGCAATGGA	AACTAATTGA	CTTTTCAATT	AACGTGATGG	ATGCTGGCGT
	20		CTTGTTTCGG	TCTATCACTT	GATGACATCT	GAAGAAGAGA	CACAACGGCA	CACCAÇGAGT	GCTAAGGGAG	AAGAAAGGAG	TCGGCGACTG	TCTGAGCAGG	ATGGACTACG	CCTTATCAAG	ATTGATAGCT
	10	Littlini	AAAAGCTCTT	TTCCTTTGTC	ATCATCTTCC	GTACGCTTCG	CGTCACACGA	GGATCTGACT	GTTGCTGGTG	TGATTGGAGG	TIGGICITIC	CATCAGCCIC	TGGTGGTCTC	GAAAGATTAT	GGTTGTGACA
		S	UBS	TIT	UTE	SH	EET	(RU	LE.	26)					

# Fig. 13. 2/2

200 260 320 500 096 ATGAGACTTG CAGTGCTCAT CCTTCTGGAA CGTTTCAGCT CTTGGGGAAA AGATGATTAT ACAAACTTTT AGAAGTTTTT CGGAAGGAG1 TCTGTGATA CTCCTCCAC AGCTAGAGT AAA 1553 AGCGAGAGAG TGTGCAGCTG GAGATCAAAC TIGGACCACG AAATGCCGCG GATTACCCCG AAAAAAAAA GTGAAGAACT ACCGGCAACT CATCCAAACC GTTATGTAAG ATAAGCTGAG TTGGATGTAT TAAATAAAA TTCTCGTGG CATCTGTGGC TCAACATCA CATCAAGACA CAATTTCACA CGAGGAATGG GCAGCGTAAC TGTCCTCGT TTCGTTATGT AGTATCTCAT TACTGGATC TTCACCTAT TTAGTGTTGG CTGGCCCATG ACGGTGTTGA TTGGTTTGTG GTCGTCATTG AGAAACTGG CCTCTTTGGA CTTCGTATCC TTATGCACAT TIGGCAAGTI ATGTATTATT CAGGACACAT AATGCAACCT GCTCAGCCAG GGTTCACAGA AATATGCTCG GGCCCCACGA TGTAAGGATG CTTTGCCTAA CACACGGAAG AGGGTATCTG GGAATATTCT TIGITGIGAT ATGGATGGGT AGAACTIGT TCCAATAAAC AGCTGTAGCG ATACTCTAAG TGCTGTGCG GAATCAATAA CGTAGGATAC GAGITGGGGT ATGTGGAATC GTCCCCTTCC GAAGAATGCG TAAAC TC TTC TTATAGCTG AACCAGAAGT GCTGTGTG

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# Fig.14. 2/2

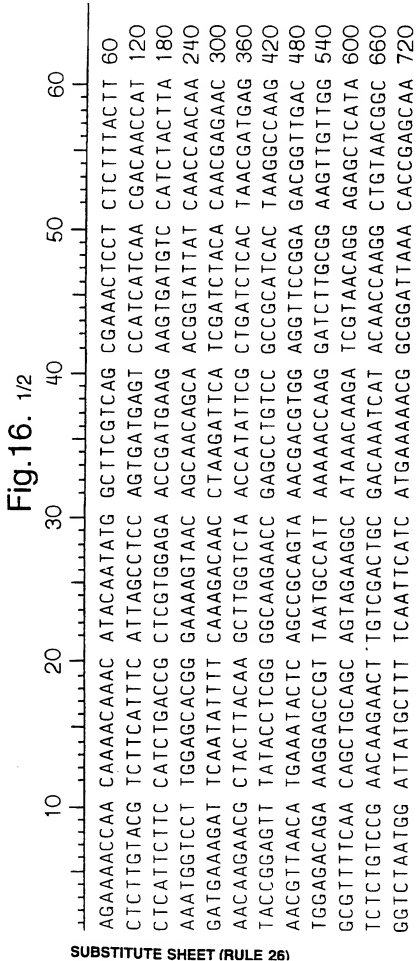
1020 1080 260 1200 900 096 GTGGCGTCCA TATTTCGAAA TAAAAAAAA TCTTTACTTA GATGAAACCG GGTGGAAGAG ATGGATCACG AGCCCAAACC AATAICATIT AACGGCGGAT GTAATCAAAT GTACGTAACT AGAAATATAA CATCATGAAA TCCTAGTAAA GGAGAGAAAC ATCTCATGCA GTATGTATAG TATGTTGGTC CAAATGCAAC GGTTAAGTAC CTATTGGATT TATTGATGCT TGGTACGAC ACGGTGTTGA AACAAAAGA ATGTATTAAG CTGGAAAGTG TATAAATCTC GAACCAATGG TGAGTGTTGC CCTCGTATCC ATTATTGTTT ACGAAGATGT ACATTAGGAT CTTTTCAATT ATGGATTATG CCTTACCACG TACCAGCCTG GGAATCTTCA GGATCAGAGA GCGATTGAAG ATTTAATCTG TGGCTTGATT GAGGATGGTT TIAATCIIGI ATCGATGGAT GTTTGAAGTT CACCATAGGG ATAAGTTAAA CGGCGGTCTA GCAAGACTAT AGCAGITICG AAGCTGGGGA AACCAGCAGT ATATITGCTT AGTIGIGACT GGTTGGTTAT GTGTGGGATT TTACCAATCT 1390 AAGGCTGTAA AATCCGGTAA CATTCAAAGA AAAAAAAAA TAAACACCGA AAAATTCGAG CGTTGAAGAG CGGTTCGTGG AAAATTGATT CITICCAACA CTGTTGTGGC CTIGGGGTAC TGGGATTGTT

Fig.15. 1/2

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		09	120	180	240	300	360	420	480	940	900	099	720	
09	Littliti	TCACCAAAAC	GAGTCCATCA	GAAGAAGTGA	AGCAACGGTA	TTCATCGATC	TTCGCTGATC	GTCCGCCGCA	GTGGAGGTTC	CAAGGAACTI	AAGATCGTAA	TCGTACAACC	AACGGCGGAT	
90	Lengther Line Hiner Lines to entering	TATGGCTTCC	CICCGGITAI	GAGAACCGAT	TAACAGCAAC	CAACCTAAGA	TCTAACCATA	AACCGAGCCT	AGTAAACGAC	CATTAAAGAC	AGGTATAAAC	CTGCGACAAA	CATAATGAAA	
04	Treatment of	AAACATACAA	TTTCGTTAGC	ACCGTTCGTG	ACGGGAAAG	TTTTCAAAGA	ACAAGCTTGG	TCGGGGCAAG	ACTCAGCCGC	CCGTTAATGC	CAGCAGTAGA	AACTIGTCGA	CTTTTCAATT	
30	Tirritini	CCAACAAAAC	TACGTCTTCG	CTTCCATCTG	TCCTTGGAGC	AGATICAATA	AACGCTACTT	AGTTTATACC	AACATGAAAT	AAGAAAGGAG	TCAACAGCTG	TCCGAACAAG	ATGGATTATG	
20	111111111	AACTAGAAAA CCAACA	ACTICICITA	CCATCTCAGT	CTTAAGATGG	ACAAGACGAA	GAACAACAAG	TGAGTACCGG	CAAGAACGII	TGACTGGAGA	TTGGGCGTTT	CGTATCTTTG	CGGCGGTCTA	
10	111111111	CAACTATCAA	TCCTCTCTTT	TCAGTGACAA	TATCCATCTA	TTATCAACCA	TACACAACGA	TCACTAACGA	TCACTAAGGC	CGGAGACGGT	GCGGAAGTIG	CAGGAGAACT	AAGGCTGTAA	

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780	840	006	096	1020	1080	1140	1200	1260	1320	1380	
TCTTTACTTA	GATGAAACCG	GGTGGAAGAG	ATGGATCACG	GTACGTAACT	GTGGCGTCTA	AGCCCAAACC	GTAATCAAAT	TATTTCGGAA	TCAAATATCA	ATAATAAAT	AAAA 1434
CAAATGCAAC	TCCTAGTAAA	TATTGATGCT	TGGTACGAAT	CTATTGGATT	GGAGAGAAAC	GGTTAAGTAC	ATCTCATGCA	GTATGTATAG	TCTAGGTTGG	TAAGAGAAAT	AAAAAAAAA
GAACCAATGG	ACGAAGATGT	TGAGTGTTGC	CTGGAAAGTG	ACGGCGTTGA	ACATTAGGAT	CCTCGTATCC	AACAAAAGA	ATTATTGTTT	TGTATAAATC	GATTATGTAT	GAAAAAAAA
CCTTACCACG	ATCGATGGAT	TACCAGCCTG	GGAATCTTCA	GGGTCAGAGA	GAGGATGGTT	GCGATAGAAG	GITIGAAGII	TTAATCTTGT	GGATTTAATC	MCTTTGACTT	TGGTTTCGCT
	AGTTGTAACT	AGCAGITICA	TTACCAATCT	GGTTGGTTAT	ACGTTGGGGA	GTGTGGGATT	AACCAGCAGT	ATAAGTTAAA	TTCACCATAG	AGAATATTTG	TCAACAGCAT
TAAACACCGA GAAAGACTAT	AGAATTCAAG	CGTTGAAGAG	CTTTCCAACA	CIGIGGIGGC	CITGGGGTAC	AATCCGGTAA	CGGTTCGTGG	TGGGATTGTT	AAAAAATGA	TTTCATTCAA	GGTATATTIC
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# Fig. 16. 2/2

1080 1140 1260 096 GAAGAGAGCA ATTGTTATAA TICTCAAAAA TCGTGGAACC AAAAAAAA TICGAGAGII CCAACATTAC TIGATICACC CAAAGAATAT CGGTAAGTGT TGTGGCGGTI GGGTACAAG TACTTAAAAA **TCGAAAAAA** AAAAAAAAA GTAACTCTTG CGTCCAAATC TCAAATTGGG CAAACCCGGT TCATTTCATT AATGATATAT ATCACGCTG1 GAAGAGCTT AAACCGCGT ATATAATAAA AAAAAAAAA AGTAAAGATG GATGCTGGTG ACGACTATGG ITGGTCAATA TGGATTGTAC AGAAACGTGG CATGCAGTAA TGCAACTCTT AAGTACAGCC GTATAGTATT CAATGGCAAA AATCTCTATG ATTAAGAGAA AAAAAAAAA TAGGATGGAG AGATGTTCCT TGTTGCTATT TGTTGACTAT GTATCCGGTT TTGTTTGTAT AAAGTGTGGT AAAAGAATCT ACCACGGAAC ATGGATACGA AGCCTGTGAG TCTTCACTGG CAGAGAACGG GAAGTTAACA TCTTGTATTA AATCTGTATA AAAAAAAAA TTGAAGCCTC TIGATTATGT ATGGTTACAT GTGACTATCG CAATCTGGAA SGTTATGGAT **IGGGGAGAGG 3GGATTGCGA** GTTAAATTAA AAAAAAAAA GITICGIACC ATAGGGATTT TIGCTITGGC GACTATCCTT AGCAGTGTTT A 1441

				Fig. 17. 1/2			
	10	20	30	01/	50	09	
SUE	Treethorn	Treeters.	1	Time the transfer of the	Linkling	List List	
BSTI	CAGAACTAGA	ACAACCAAGC	CAAACATACA	ATATGGCTTC	CTCAACAAAA	CICATITCII	09
TUT	TACTTCTCCT	ATACGTCGTC	GTTTCATTAG	CCTCCGGTGA	TGAGTCCACT	ACCATTAACA	120
E SI	ACCATCTCAA	TCTTCCATCG	GACGGCTCAT	GGAGAACCGA	TGAAGAAGTG	AGGICCAICI	180
HEE	ACTTACAGTG	GTGTGCGGAG	CACGGGAAAA	CTAGCAACAA	CAACGGTATC	GTCAACCAAC	240
T (R	AAGACGAAAA	GTTCAATATT	TTCAAAGACA	ACCTAAGGTT	CATTGATCTA	CACAATGAGA	300
ULE	ACAACAAGAA	CGCTACTTAC	AAGCTTGGTC	TCACCATATT	CICIGATCIC	ACTAACGATG	360
26	AGTACCGGAG	GTTATACCTC	GGGGCAAGAA	CCGAGTCTGT	CCGCCGCATC	ACTAAGGCCA	420
)	AGAACGTTAA	CATGAAATAC	TCGGCCGCAG	TAAACGACGT	GGAGGTTCCG	GAGACGGITG	480
	ATTGGAGACG	GAAAGGAGCC	GTTAATGCCA	TTAAAAACCA	AGGAACTIGC	GGAAGTTGTT	940
	GGGCGTTTTC	GACAGCTGCA	GCAGTAGAAG	GTATAAACAA	GATCGTAACA	GGAGAACTCA	009
	TATCTCTGTC	CGAACAAGAA	CTTGTCGACT	GCGACAGATC	CTACAACCAA	GGCTGCAACG	099
	GIGGITIAAI	GGACTATGCT	TTTCAATTCA	TCATGAAAAA	CGGCGGTTTG	AACACCGAGC	720

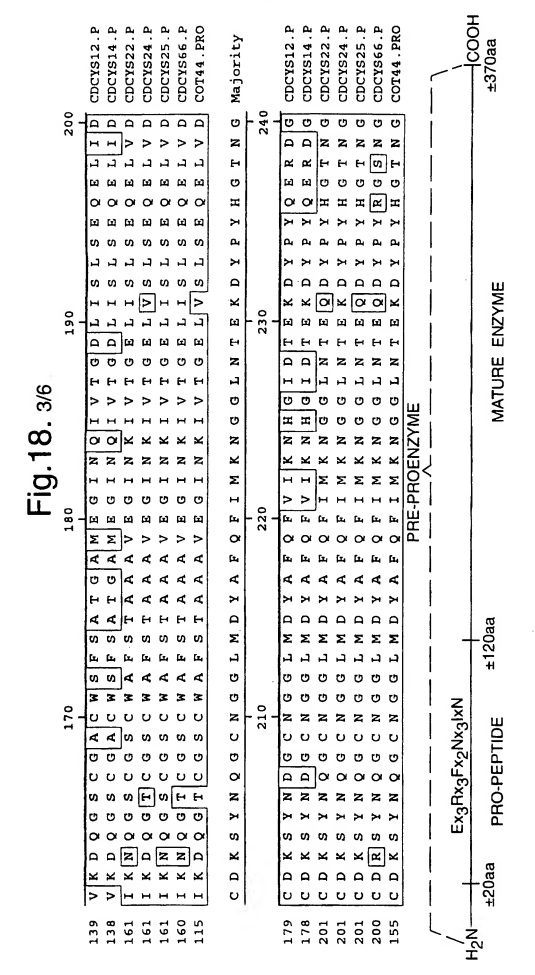
# Fig. 17. 2/2

260 780 AAGTCCGGCA CTCTTAAAA AATTCAAGAG TGAAGAGAG CCGGTTCGTG TGATCCAGC CAAAGAAAGA GTGGTGGCTG GGGGTACAC **TAACTAAGAT** AAAAAAAA TCCAACAT TITACIGAAG AAGGAACTCG AAAAAAAAA **IGAAACGGCG** AGATCATGCA GGCAAGGTCC CAGTCCAAAC GGGAGTTTTA AAGAAGGTAT TCATTTCGTT TGATATATT **IGGAAGAGTT** TCGATCAATA AAAAAAAAA AATGCAATTC CTACTGAAGA TTGAAGCTGG CGGTTAAGTA TATATCAAAA TATAATAAAA AAACTCAATT GGACAAATCT AGAGAAACTT ATTGGATTG1 **AAAA 1474** TCCAATGGAA AAAAAAAAA AGTGTTGCCA GGAAGTGTG ATTAGGATGG TTTTAAAATA GTATGTATAG AATCCTTATG TTAAGAGAAG AAAAAAAA GAAGATGTTC GGTATTGACT GCCTCGTACC AAAAAAAAA AAAAAAAAA TTACCGTGGT TGATGGTTAC CCAGCCCGTG GATCTTCACT TICAGAGAAC GGATGGTTAC TGCGGTTGAA TGTTTGAAGT ATTATTGTTT AGTCTGTATA TGTTTATGTA TIGGITATEG GITGGGGAGA AAAAAAAAA AAGATTATCC TTGTAACTAT ACCAATCGGG CAGITICATA AGTGTGGAAT SAACCAGCAG TTAATCTCAT TIGATITGGT AAAAAAAAA ATACGGATTT

Fig. 18. 1/6

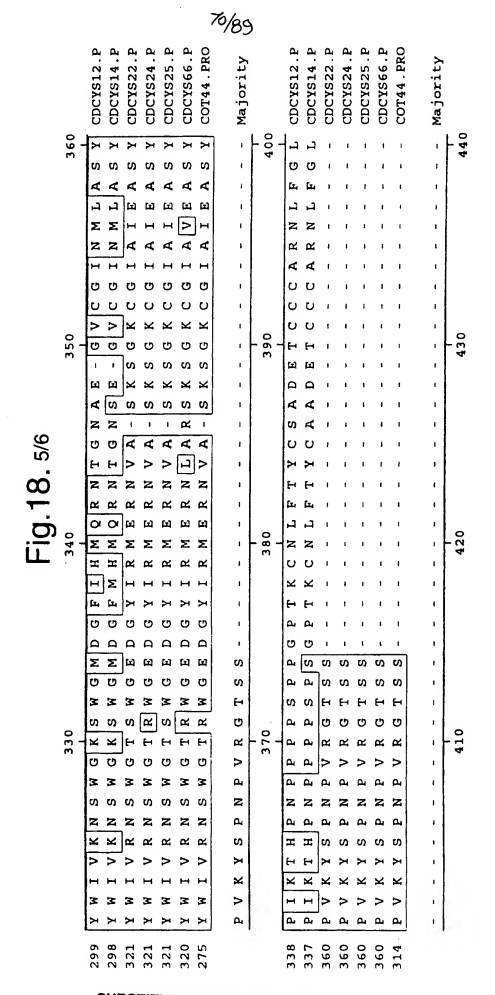
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Decoration 'Decoration #1': Box residues that match the Consensus exactly.

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Fig.19. 1/2

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Fig. 19.

# 73/89

1020 140 380 560 080 200 260 440 900 096 999 840 780 TTTCTAAAG ATATTGTGAC ATATTCAAAC ATAAAATGTC TGTTATTATA AATGTATTTG ACTAATTGAA **FCCAACAAGG 3GCCCAATCA** TCAGITAAAC TCTTGGTTAC AATGATTTAT AAACGCTTAT TCTTTCAAC T C T T G T T T T **AAGTAGCCT**1 CATCTTAGG CAAAGTAT AAAGICICC AATACATATA ATGTCACCGA ATATAACCTG TACTTACAAA TATTCTCTAA AAGATTGGTA CGTAAGTAGG TTTTGCTTA TAGTAGTAAA CTATCATCAT AAATACTATT ACTATCTTCT AAAGCATATT TATTATTAT AAAACAAAC **ACAAAAGCTC** CITACIGGCI AGCCIGTC ATTCAACGC TTTAGCCAGG **TAAAGTAGAG** AGCCTACCTA AGITGIAGA AGAATTGGAA TTATATACAG CGACGACCAA **ACATGGGACA** TGTTGACCAA SCCACGCCAC TCGTTCTATT ATACACTGGA CGTGATCTTA GCCATAAAGT AGTAGCTTTC SCCCTTCAGC TICTAAACT STAATGGGCC TIGITGCAT CTTTATACTG TAAAATTTAA AATCCACGA IC AAC TC TAG **ITGCTGATTA** GATGTAGTAA CTGGTCTCCG AAGGGTTAAT AAAGGTTGAA TAATAAAA TTGTAAACG **AAGACTGAAC** AAACCTTTTT AAATGTTCTT **ICTTTGTCAC** TTTCTAAAGT ATGAATTTAT ACATGCATTT AATTTIGATT AATG 1644 AGAGAAATTA AGTAAACTAA ATATACTAAA TAACCTAAA ACATTTAACA GTAGCATTTG **IGTTAATTAC** AAACAACAAT CAGITAGGIA GGTCTAATCA TAAGAAAGA AGCAGTTTTC CGACCTTATT ACGITCGIAI TGTGTTTGTT **IGTGATTAAT** CTACTACTCT ACTTATGTTT AGCCGCGCGT ACAAACTATC ATCGATTTCA ATTAATAAGA TTTGATCTA CCTTCTGATA TACTIGICGA AGTGTGAAGA CACTAGCCAC ATATATATA TACTAGGTA AGTGGTCTTA TAGCTAAGA **TATTACTAGG** TTATGTATTT AATAGTGACC AGAAAAGGC ATTAAAAGC GCTATGTTT TCATAACCG GGTTTCCCTG ACAAACGCT

Fig.20A. 1/3

3997 bases 452 bases 21697 bases 28 sequences 20 sequences	Contig 5: Contig Length: Average Length/Sequence: Total Sequence Length: Top Strand: Bottom Strand: Total:
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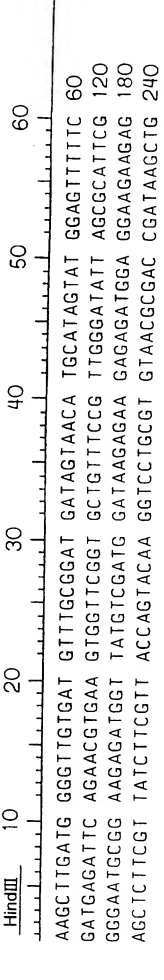


Fig.20A. 2/3

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AGAGTGCGGT	GGTTTGTATT	AGGAGTTTGG	ATCCTTCGAT	GTGTCGAAGC	AGATGTTAAC	TTTTTGTTAT	AAACGAACGC	TCATTATCCC	CAACAGCAAC	GAAGCTAAAC	AGAAACTGAG	AAGITITATA	AGAATCAACG	AGCAAAAGGG	CTTTCTTTTG
TACGCGGGGA	GAGTGGTGTG	GAGTGGGTTA	CTGTTGAATG	TGGTGGCTTC	TTTGCTCTCA	TTTGGTTCTG	AGTGGCGGCT	AGATCAAATG	GCAGCAGCAG	AAATAAACAC	TCATCAAGAA	TIGITIGCCA	GCGAATAGTA	GAGCCCCCAC	TTCGATATAT
TACGTTGCCA	GCAGAAGCTG	GAGTAAGCCG	TCCTTTGTCT	GGTTCTGCTT	TATGCCTTCT	GTTAGTGATG	TAGAAATCAC	CTAAAAAGAA	AGCACCAGCA	GAAGTAAGAG	CCACTAAACA	TATCATTCAT	TCCAGATACC	TATGATGATG	TTCGTTTATT
ATGTTTTCAA	CGGTTTTGCC	ATGAGGCTGA	<b>gCGTGGAGAG</b>	GTGGAAGACA	CICCGGGTA	CTAGGAGTTA	GATCCTCCGT	GAGCCACCTA	GTGATACCGA	CCGAAACGCA	GITICITCII	ACTATTTTAT	CTTATTCTCC	GCTTAGATTT	TTGATGATCA
ATCGGTTGGC	GATGTAGCTG	ATTCTTTGGG	GAGAACGAAG	CITGGAAGCT	AGCAAGTTCC	AATGTTCAAA	ATGGGTGATT	TGTTCCACCT	CAACGIGGIG	AGTGCGGAAA	GACAACACAA	GATCATITCT	AAGAGATCAT	AAGAACAAAG	TTTAGGGGAA

Fig.20A. 3/3

## 76/89

### 2040 2100 1740 1860 1980 GGAGGATTGG TATAAACAGG TITIAGAGAG CTTCATAAGG GAAGACATIG TAGAGTAAAC AATCAATCCG GITGCAAAAA TTCTCAATC CTATTTCTT AATCAAATTT AAACGAAAAC CTATTAGATT AAGCCTCAA GAATCATTC TGTTCTTGAA TGTAAGCATA ATACAAACA TTACTTATAA TAAAAGTTCA AACTATTAG **TATTGGAGAT** CTGAAATAAT CTTTGACCAC TITCAACCT AGCAGGTGAC AATATGAGGT TCAGTTGCTT TTTATTATT TCCCTCAC **TAGITICITI** TCTAAAGAAA AAGAAAGTAA AGATTCATAA **TAGGTATGTA** TITIATIACG CCAAAATTTA ACTATGGACC CAGTATTAGT ATTTATATGT TTTGTTTCAT AATTACACTT TCATTTCTTT TTTACTACTT TTAACCACTI CAAAGAATTA TTGGGTTGTA CATTAGTGAA TTCTTTGGAA AAGAATTGTG **ICATAGCCTA** AATTATGGGA ATTAATCCAC GACTGAAAAC AAAATATCCT CACAGAAATT GAATTTGTC TATAGTATTT CATGTCTTCT GITITIGCI GTATTAGTAA TATGTATTTA CTTCGTAAAA **TGAGACAAA** AAATTAACGA AACAAAACT ITATCCCCCA CAGAACATCA TACAC TGCGG ATGTTAAACC **TAATAGTATG** ATTTTAACA GGTTTCTTCT TATTTAGAT GAGTTCGTTT AAGGCTTAGA GAACTTATCT AAACAATACA **TTGTGAAAG** ATTGAGTTAG GATAGAAGTG ATATTATGCG T AC T C C T G C GAAAGATAC **LITTICTIAG** GAACACAAG CATTTCTGTT GCGGTTTTGT CAAACATATC CGGATTAGGT

Fig.20B. 1/2

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40 2150 2	ACGATATCAA	CAGAAACACG	<b>IGAAACITGG</b>	TTGTTAAAAT	ATGGGGCACC	TICATICIAG	AGAGCTCTGT	AATAGAAACT	ATTCCTCATA	TICIAGIGGI	ATATGTATCT	CTTTTCGAG	ATTATTGGAA
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2130	CTTCGT	AACGGT	CCTTAT	TITCCT	AGAAAG	TGAAAC	TTAATA	CAACTA	ATG	ACTCAC	GTATTT	TICCCA	GTTTTT
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<u> </u>	616	AAT	116	CTC	AG I	AAG	ATG	ATT	TAC	ATA	AAI	AAT	CTC
	STCG	ACCG	3C AC	AC AA	ATAA	r G A C	TAT	ATTT	<b>JEAC</b>	AAA	ATAG	AAAA	CAA
21   1	CAAACAGTCG	AACCAAACCG	AGCGAGGCAC	TTAAAGACAA	GTCTTCATAA	GCCTCGTGAC	ATTCTTCTAT	ACGTATATTT	ATATTCAGA	TTTTGATAAA	CGATATATAG	TTAAACAAAA	GTAGTTGCAA
	CAA	AAC	AGC	TTA	GTC	ງງ၅	ATT	AC G	ATA	111	CGA	TIA	GIA

3900

AAATACCTAA

TGAAATATTA

AATCAAGCCA

TCCCACCAAA

CACATTCTTG

T A A A T A G T T T A A A C C A A C A A

CCATTATTAC

TCTTGCATGC ACTTTGGTTA CAAAACTAGA

TITITATIAT

AACAAACATA

AATACCAACT

3960

TACCAACTAT

# Fig.20B. 2/2

3360 3480 3240 3300 3540 3840 3600 3660 TTCCATGAGA ATTICGTTAA GCCAAAAAA AAGTCAAAAA TAGATCAGCG TAACCCAAAA TACTAAATAC AATGATCTTC TATATATT AACAACAAAC GAAGATATTA ACGICCACIA TTATACGGCT TCAGTGAATC CTCCAAATT ATC TTTGAG GTGGTCGGTA **IGACGTTATA** CAAATATGAG ACGCTCTGTG TGTTGATGTA TCCATTIGAA CTATACCTAT CITIACAACI GAAAGATTT TACGAGATAA AACCCCGTAG CTTTTTAGA AGATGCAGCT TAATAATTAT TGTTATTATT AATTAAAGT GTGGATTTAA CACACGAAAA TCAAAAATAA TCACAGACAT AAAGAATGG TACGTACATC GAATGGTTTG TCGAAATGCT TATATACTAT CTATTTACGT AGTCGATAGA CCGAATAAAA GCAAAAAAA CTGATGATAT ATGTTGTATT AATATCTAGT GAAATTCACA ATGCATGTCT AGAAATTAAT TATATGGTAG TCATGCTGCT TTTGTTTGC ATCTCTCGTG AAAAAATATA TGATACTACG ATCAAACATA TIGAACGACT TCTGACCGCT CGAAGAAAAT AATTACAGAC ATCCGATTGT CCATATTTCG CGTAGATTCA TTCGACGTAA AATTGTCACA **ICAAGTGACA** AAGCGAATTG GTATGCAGGA AAATAACTGC CATACTTACA GTGTTCGGAA ATAAACGTAA AAACTCTTTG TTAAGAGCA GATTTCTAAA ACAAGATGGT CCTATTTACC AAACAGTTGT ATTGATGTAA CICAAACCAI TCCAATTAGA ACGCGGTTCA AAGITGAICC ATGTTCCCAT TAAACCGCTT TGATCAAAC CACGGAAAAT ATTGTATTAT AAAGAATAGT ITGAATTGTA AATAAATTTA ATCGTCTACT TATATTAGE STAGITITIT

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CAATATG START

3997

Fig.21. 1/3

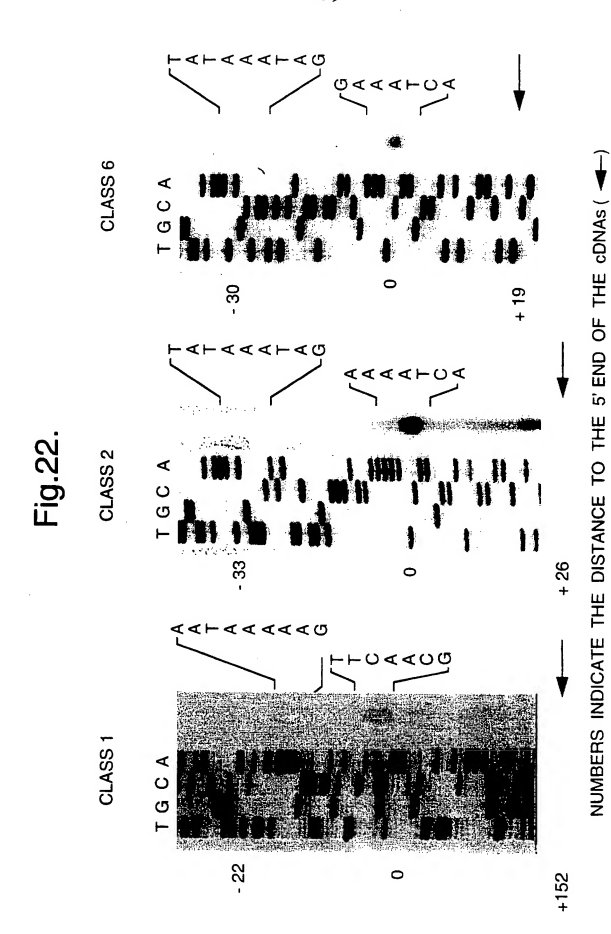
Contig 2: Contig Length: Average Length/Sequence: Total Sequence Length: Top Strand: Bottom Strand: Total:	0 4 60	2000 bases 413 bases 9920 bases 14 sequences 10 sequences 24 sequences	nces nces		
BamHI 10 20	30	40	50	60	
GGATCCCACA CATAACIGTA ATGTTTCAAT ATTTAGTAAA ATCCGCATTC TGGCCCAATA ATGTATCTAG GATTTTTATA GTACTATGTC AGCTTATCAA AAAAAATTCT GAAACTTTTC		ACTCACGTGT AAACTTATAA TCCACCGATA ATTTTCACCT	AACTTTGATC GTGGGCTAAA AACCGAAGCG CTCTATATCT	ATCGAAACCT 60 TCTCTTTTGT 120 TTACCCTTTT 180 CCAACGATCA 240	

# Fig.21. 2/3

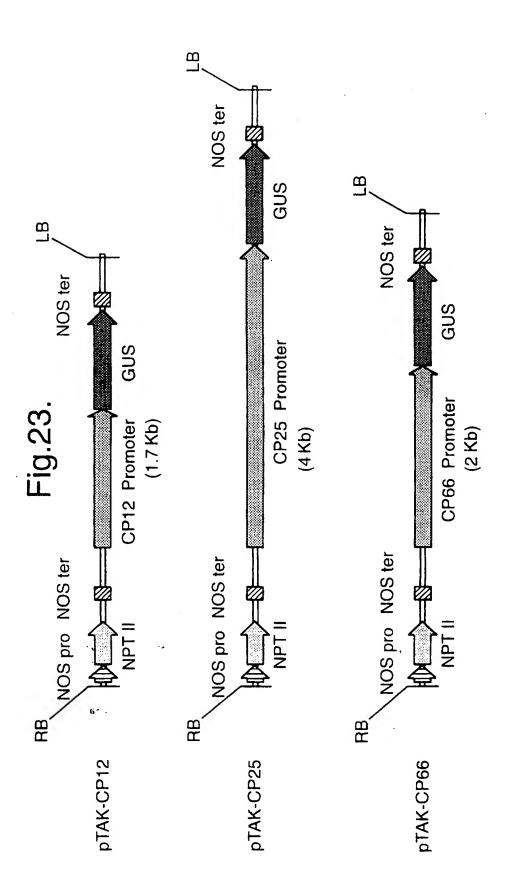
080 999 600 720 480 540 780 840 096 006 CCTCAAACAA TTGTGAATCA **ACTITCTATG** ATACTICAAG AAGTTATTG TGAAGGACGG CTGTTCATTG GGCTTTTAGC TGTCCCCGCC GGTACCACCA TITIGCIGAT CTTTGAAGTT AACTTAAGCC GCGATGATTC TIGGITIGAT GTGGTGCAA ATCAGTTTTA ATTTCAACA ATATATATT GAATTGTCGC TGACAATTTG CAGGCGGAGC CGTGCTATGA AGTCTACAAA GATTTCCTCA AATCGGTCGT ATCCCAGTTC TGATTGATCG CTATAACGGA TTGTATGTTA AGATTGAAAC TAACACCGGT TTCTGATGAG GTCGTGCCAA GTTCGGCTCA TAGATTTACG CCGICTCAGI GGTTCAGTTG CTCCTAAATT GAAAGGTAT CCACGGTTCT TIGTGAAAGT TGGTACGTCT GTCGGCTGTC TAATGITATT TGAGTTTCTT GTTGTTGCT TATTCAAGC TATTAAATTA TAGTACCCAG CTGAAACGAT GITTAAICAA CTCCAGTGTT AGGGGGTCGA GTAAACTCTG TTGTGGTGTG AAGGTGTGCA GAATCTCTGG ACTTCGAAAC CTGCAAGGCT GCATGACAAT ATTCTGAGAT TCTGGTTTTT GGCTATGTAT CGITICACCI CCATATCTCT AGATGTCAGC GCCTAGAACA TCTGGGCAGA CTAACAGAGG GCGCCTATTG ATGGTAACCA TACCATATIT GATGTGATCG TCGCTACGAT CTAATATGAT AATCCAGATC TTTTTTATGT ITTGIGGAT GAGAAAAAT ACCTIGAAAC GAATACTATG CTACCATCCA GTTATGGTAC CAACCTICGA GTCCGATTAA ACCTTTCTTC CGTATGTTTC GTTGCTATGA TACAACTAGA TTCCTTGGAG GAAATAATGA TACATTGGCA **IGATGTAAAT** CCCTAACATG GAGGATGGAC

Fig.21. 3/3

1260	1320	1380	1440	1500	1560	1620	1680	1740	1800	1860	1920	1980		
AAATTTATTG	TCGAGGGACA	TCTGGCTTGG	CTAGGTCGAT	CAAGATCCAG	CCGTACCTTC	CGTCTAATTA	TAGTTGATAT	CTCCCTGCCT	ATTAATTTCA	ATTACTGCAT	TTTGGTTATA	CTAGAACAAC		
TAAAAACTTG	GTGAACTGCA	AACGAGTGTG	TGCAGGCTCT	AGCTACTTGC	ATAGCAATCA	ATGGAAAAAT	GTTATCTTGA	ACATAAATAT	AAGGTGTAAA	TIGCATGCCC	ACTCCAAGAA	ACTATCAGAA		
TCTTGAGACT	ACCTGCTGCG	AGGTGTTGAA	TCTATGTTGT	ATGTATTCAT	GAAGCCATTA	ATAGAGAACT	CGTATTTATT	ATTAGAAAAT	GAAGATATTT	GTCCACCATC	AAATATTAAA	AGCCATATCA		
CATAGTATCA	ACATTAATAA	ATAGGGAATA	AACCICACCG	ATAGIGITIG	CAAATTGATG	AGATACCATC	ATCAGCAAAC	TICIGAAATI	CGTGGGGAAT	CATTTGAACC	TACCAACTIG	AACGAAATCA	2000	
TGTCTTTAG	TTATTCTCCG	TAGAAGAGAG	TATACTCAAG	TGCCAGCAAC	GGAGAGCICI	CCATCACTGA	TATATTATAG	TAACTAAATT	AACGTTCTTA	CGGCAATATC	TAGAAAGTA	CTCAGCCTCC	ATACAATATG	START
TTTAAAACAA	GICAGATAAC	TGGCAGTAAT	TGGAGCTCAA	TCAAATCTCA	CCCAAAACAT	ACTATCTCCA	TACTIGGGTA	AGTATATAAG	ATTATCACAC	TICATATITC	TITATTAAA	AATAGCTCCT	CAAGCCAAAC	
	TGICITITAG CATAGIAICA TCITGAGACI TAAAAACIIG AAAITIAIIG 1	TGICTITIAG CATAGIAICA TCITGAGACI TAAAAACITG AAAITIATIG 1 TIAITCICCG ACATIAATAA ACCIGCIGCG GIGAACIGCA ICGAGGGACA 1	TGICTITIAG CATAGIAICA TCITGAGACI TAAAAACITG AAAITIATIG 1 TTATICICCG ACATTAATAA ACCTGCTGCG GIGAACTGCA TCGAGGGACA 1 TAGAAGAGAG ATAGGGAATA AGGTGTIGAA AACGAGIGTG TCTGGCTIGG 1	TGICTITIAG CATAGIAICA TCITGAGACI TAAAAACTIG AAATITATIG 1 TTATICICCG ACATTAATAA ACCTGCIGCG GIGAACTGCA TCGAGGGACA 1 TAGAAGAGAG ATAGGGAATA AGGIGIIGAA AACGAGIGIG TCTGGCTIGG 1 TATACICAAG AACCICACCG ICTAIGIIGI IGCAGGCICI CTAGGICGAI 1	TGICTITIAG CATAGIAICA TCITGAGACI TAAAAACTIG AAATITATIG 1 TTATTCICCG ACATTAATAA ACCTGCTGCG GTGAACTGCA TCGAGGGACA 1 TAGAAGAGAG ATAGGGAATA AGGTGTTGAA AACGAGTGTG TCTGGCTTGG 1 TATACTCAAG AACCTCACCG TCTATGTTGT TGCAGGCTCT CTAGGTCGAT 1 TGCCAGCAAC ATAGTGTTTG ATGTATTCAT AGCTACTTGC CAAGATCCAG 1	TGICTITIAG CATAGIAICA ICITGAGACI TAAAAACITG AAATITATIG I TTATICICCG ACATTAATAA ACCTGCTGCG GTGAACTGCA TCGAGGGACA 1 TAGAAGAGAG ATAGGGAATA AGGTGTTGAA AACGAGTGTG TCTGGCTTGG 1 TATACTCAAG AACCTCACCG TCTATGTTGT TGCAGGCTCT CTAGGTCGAT 1 TGCCAGCAAC ATAGTGTTTG ATGTATTCAT AGCTACTTGC CAAGATCCAG 1 GGAGAGCTCT CAAATTGATG GAAGCCATTA ATAGCAATCA CCGTACCTTC 1	TGTCTTTTAG CATAGTATCA TCTTGAGACT TAAAAACTTG AAATTTATTG 1 TTATTCTCCG ACATTAATAA ACCTGCTGCG GTGAACTGCA TCGAGGGACA 1 TAGAAGAGAG ATAGGGAATA AGGTGTTGAA AACGAGTGTG TCTGGCTTGG 1 TATACTCAAG AACCTCACCG TCTATGTTGT TGCAGGCTCT CTAGGTCGAT 1 TGCCAGCAAC ATAGTGTTTG ATGTATTCAT AGCTACTTGC CAAGATCCAG 1 GGAGAGCTCT CAAATTGATG GAAGCCATTA ATAGCAATCA CCGTACCTTC 1 CCATCACTGA AGATACCATC ATAGAGAACT ATGGAAAAT CGTCTAATTA 1	TGTCTTTTAG CATAGTATCA TCTTGAGACT TAAAAACTTG AAATTTATTG TTATTCTCCG ACATTAATAA ACCTGCTGCG GTGAACTGCA TCGAGGGACA TAGAAGAGAG ATAGGGAATA AGGTGTTGAA AACGAGTGTG TCTGGCTTGG TATACTCAAG AACCTCACCG TCTATGTTGT TGCAGGCTCT CTAGGTCGAT TGCCAGCAAC ATAGTGTTTG ATGTATTCAT AGCTACTTGC CAAGATCCAG GGAGAGCTCT CAAATTGATG GAAGCCATTA ATAGCAATCA CCGTACCTTC CCATCACTGA AGATACCATC ATAGAGAACT ATGGAAAAT CGTCTAATTA TATATTATAG ATCAGCAAAC CGTATTTATT GTTATCTTGA TAGTTGATAT	TGICTITIAG CATAGTATCA ICTIGAGACT TAAAAACTIG AAATTTATIG TTATICICCG ACATTAATAA ACCTGCTGCG GTGAACTGCA TCGAGGGACA TAGAAGAGAG ATAGGGAATA AGGTGTTGAA AACGAGTGTG TCTGGCTTGG TATACTCAAG AACCTCACCG TCTATGTTGT TGCAGGCTCT CTAGGTCGAT TGCCAGCAAC ATAGTGTTTG ATGTATTCAT AGCTACTTGC CAAGATCCAG GGAGAGCTCT CAAATTGATG GAAGCCATTA ATAGCAATCA CCGTACCTTC CCATCACTGA AGATACCATC ATAGAGAACT ATGGAAAAT CGTCTAATTA TATATTATAG ATCAGCAAAC CGTATTTATT GTTATCTTGA TAGTTGATAT TAACTAAATT TTCTGAAATT ATTAGAAAAT ACATAAATAT CTCCCTGCCT	TGTCTTTTAG CATAGTATCA TCTTGAGACT TAAAAACTTG AAATTTATTG I TTATTCTCCG ACATTAATAA ACCTGCTGCG GTGAACTGCA TCGAGGGACA 1 TAGAAGAGAG ATAGGGAATA AGGTGTTGAA AACGAGTGTG TCTGGCTTGG I TATACTCAAG AACCTCACCG TCTATGTTGT TGCAGGCTCT CTAGGTCGAT 1 TGCCAGCAAC ATAGTGTTTG ATGTATTCAT AGCTACTTGC CAAGATCCAG I GGAGAGCTCT CAAATTGATG GAAGCCATTA ATAGCAATCA CCGTACCTTC I TATATTATAG AGATACCATC ATAGAGAACT ATGGAAAAT CGTCTAATTA I TATATTATAG ATCAGCAAAC CGTATTTATT GTTATCTTGA TAGTTGATAT I TAACTAAATT TICTGAAATT ATTAGAAAAT ACATAAATAT CTCCCTGCCT I AACGTTCTTA CGTGGGGAAT GAAGATATTT AAGGTGTAAA ATTAATTTCA I	TGICITITAG CATAGIATCA TCTTGAGACT TAAAAACTTG AAATTTATTG 1 TTATTCTCCG ACATTAATAA ACCTGCTGCG GTGAACTGCA TCGAGGGACA 1 TAGAAGAGAG ATAGGGAATA AGGTGTTGAA AACGAGTGTG TCTGGCTTGG 1 TAGAAGAGAG ATAGGGAATA AGGTGTTGAT TGCAGGCTCT CTAGGTCGAT 1 TGCCAGCAAC ATAGTGTTTG ATGTATTCAT AGCTACTTGC CAAGATCCAG 1 GGAGAGCTCT CAAATTGATG GAAGCCATTA ATAGCAATCA CCGTACCTTC 1 TATATTATAG AGATACCATC ATAGAGAACT ATGGAAAAT CGTCTAATTA 1 TAACTAAATT TTCTGAAATT ATTAGAAAAT ACATAAATAT CTCCTGCCT 1 TAACTAAATT TTCTGAAATT ATTAGAAAAT ACATAAATAT CTCCCTGCCT 1 CGGCAATATC CATTTGAACC GTCCACCATC TTGCATGAA ATTAATTTCA 1	TGTCTITTAG CATAGTATCA TCTTGAGACT TAAAAACTTG AAATTTATTG 1 TTATTCTCCG ACATTAATAA ACCTGCTGCG GTGAACTGCA TCGAGGGACA 1 TAGAAGAGAG ATAGGGAATA AGGTGTTGAA AACGAGTGTG TCTGGCTTGG 1 TATACTCAAG AACCTCACCG TCTATGTTGT TGCAGGCTCT CTAGGTCGAT 1 TGCCAGCAAC ATAGTGTTTG ATGTATTCAT AGCTACTTGC CAAGATCCAG 1 GGAGAGCTCT CAAATTGATG GAAGCCATTA ATAGCAATCA CCGTACCTTC 1 CCATCACTGA AGATACCATC ATAGAGAACT ATGGAAAAT CGTCTAATTA 1 TAACTAAAATT TTCTGAAATT ATTAGAAAAT ACATAAATAT CTCCCTGCCT 1 AACCTAAATT CGTGGGGAAT GAAGATATTT AAGGTGTAAA ATTAATTTCA 1 CGGCAATATC CATTTGAACC GTCCACCATC TTGCATGCCAT 1 TAGAAAAGTA TACCAACTTG AAATATTAAA ACCTTAAATAT CTCCTGCCT 1 AGCAAAATT TTGAACC GTCCACCATC TTGCATGCCAT 1 TAGAAAAGTA TACCAACTTG AAATATTAAA ACCCAAGAAA TTTGGTTATA 1	TGTCTITTAG CATAGTATCA TCTTGAGACT TAAAAACTTG AAATTTATTG 12/ TTATTCTCCG ACATTAATAA ACCTGCTGCG GTGAACTGCA TCGAGGGACA 13/ TAGAAGAGAG ATAGGGAATA AGGTGTTGAA AACGAGTGTG TCTGGCTTGG 13/ TATACTCAAG AACCTCACCG TCTATGTTGT TGCAGGCTCT CTAGGTCGAT 14/ TGCCAGCAAC ATAGTGTTTG ATGTATTCAT AGCTACTTGC CAAGATCCAG 15/ GGAGAGCTCT CAAATTGATG GAAGCCATTA ATAGCAATCA CCGTACCTTC 15/ CCATCACTGA AGATACCATC ATAGAGAACT ATGGAAAAT CGTCTAATTA 16/ TATATTATAG ATCAGCAAAC CGTATTTATT GTTATCTTGA TAGTTGATAT 16/ TAACTAAATT TTCTGAAATT ATTAGAAAAT ACATAAATAT CTCCCTGCCT 17/ AACGTTCTTA CGTGGGGAAT GAAGATATTT AAGGTGTAAA ATTAATTTCA 18/ CGGCAATATC CATTTGAACC GTCCACCATC TTGCATGCAT 18/ CGGCAATATC CATTTGAACT GAAGATATTAAA ACTCCAAGAA TTTGGTTATA 19/ TAGAAAAGTA TACCAACTTG AAATATTAAA ACTCCAAGAA TTTGGTTATA 19/ CTCAGCCTCC AACGAATCA AGCCATATCA ACTATCAGAA CTAGAACAAC 19/	TGTCTTTTAG CATAGTATCA TCTTGAGACT TAAAAACTTG AAATTTATTG 128  TTATTCTCCG ACATTAATAA ACCTGCTGCG GTGAACTGCA TCGAGGGACA 138  TAGAAGAGAG ATAGGGAATA AGGTGTTGAA AACGAGTGTG TCTGGCTTGG 138  TATACTCAAG AACCTCACCG TCTATGTTGT TGCAGGCTCT CTAGGTCGAT 149  TGCCAGCAAC ATAGTGTTG ATGTATTCAT AGCTACTTGC CAAGATCCAG 159  GGAGAGCTCT CAAATTGATG GAAGCCATTA ATAGCAAAAA CGTACCTTC 159  CCATCACTGA AGATACCATC ATAGAGAACT ATGGAAAAAT CGTCTAATTA 169  TATATTATAG ATCAGCAAAC CGTATTTATT GTTATCTTGA TAGTTGATAT 169  TAACTAAAATT TTCTGAAATT ATTAGAAAAT ACATAAATAT CTCCCTGCCT 179  AACGTTCTTA CGTGGGGAAT GAAGATATTT AAGGTGTAAA ATTAATTTCA 189  CGGCAATATC CATTTGAACC GTCCACCATC TTGCATGCCAT 189  CGCCAATATC CATTTGAACC GTCCACCATC TTGCATGCAT 189  CGCCAACTCC AACGAAATCA AGCCATATTAAA ACTCCAAGAA TTTGGTTATA 199  CTCAGCCTCC AACGAAATCA 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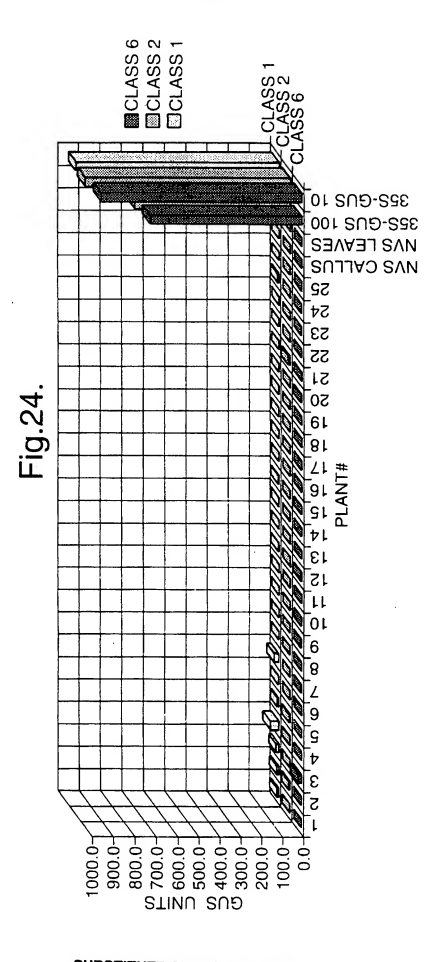


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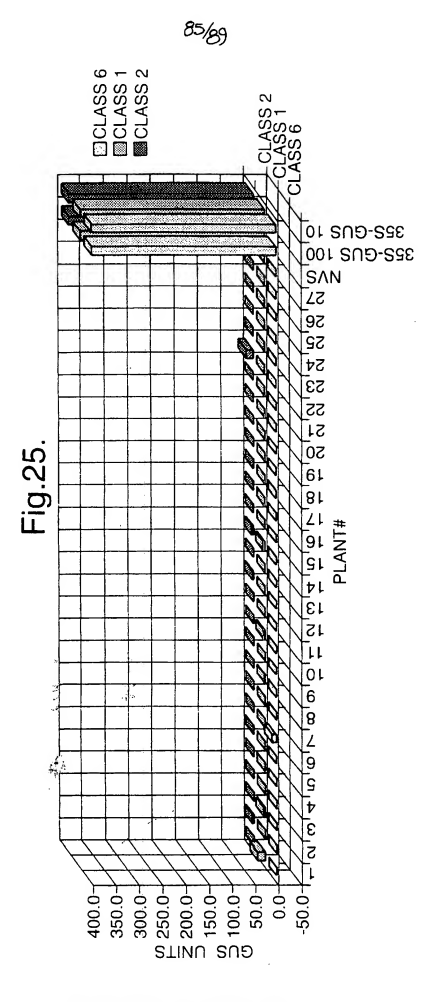


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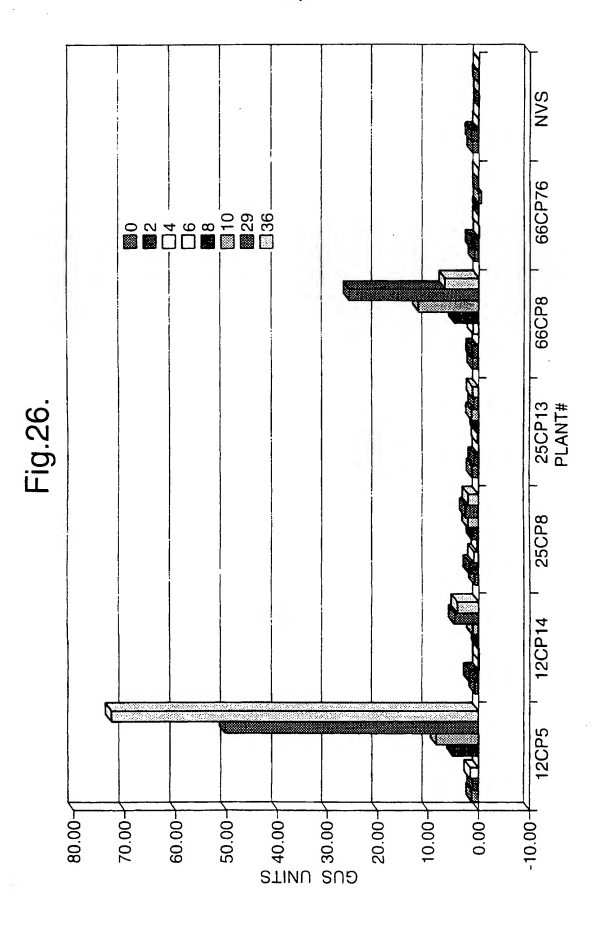


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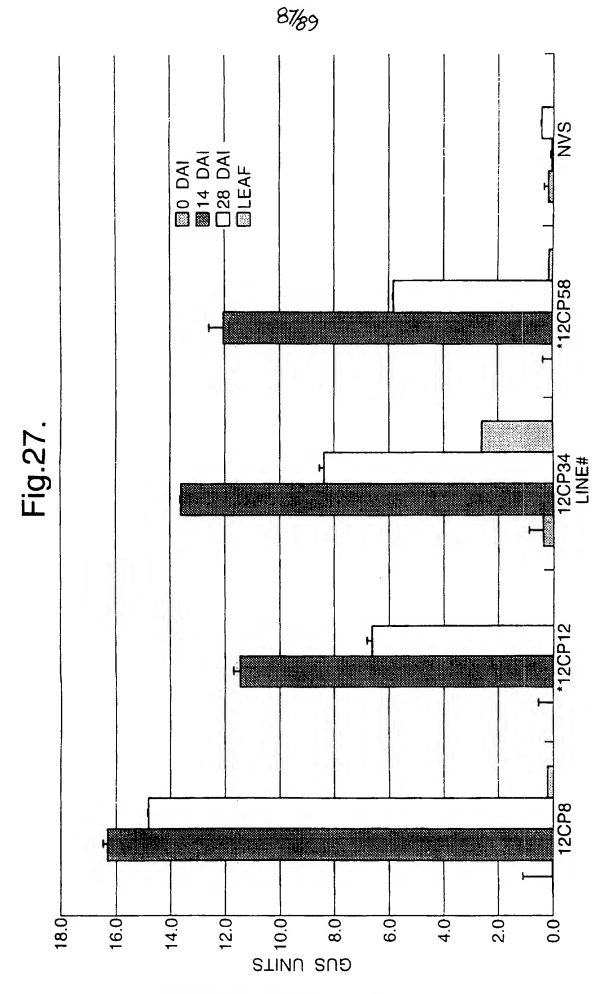


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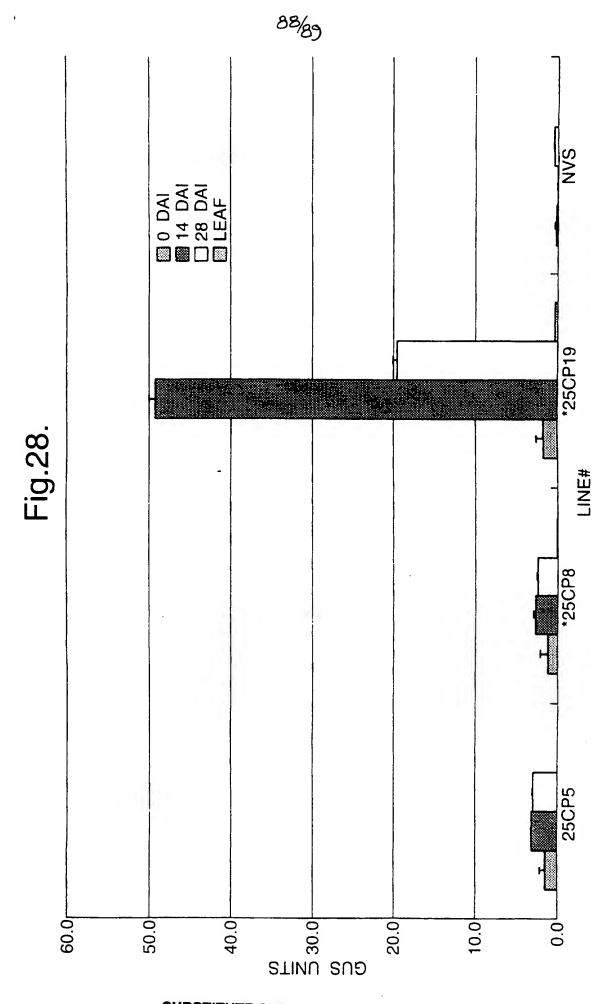


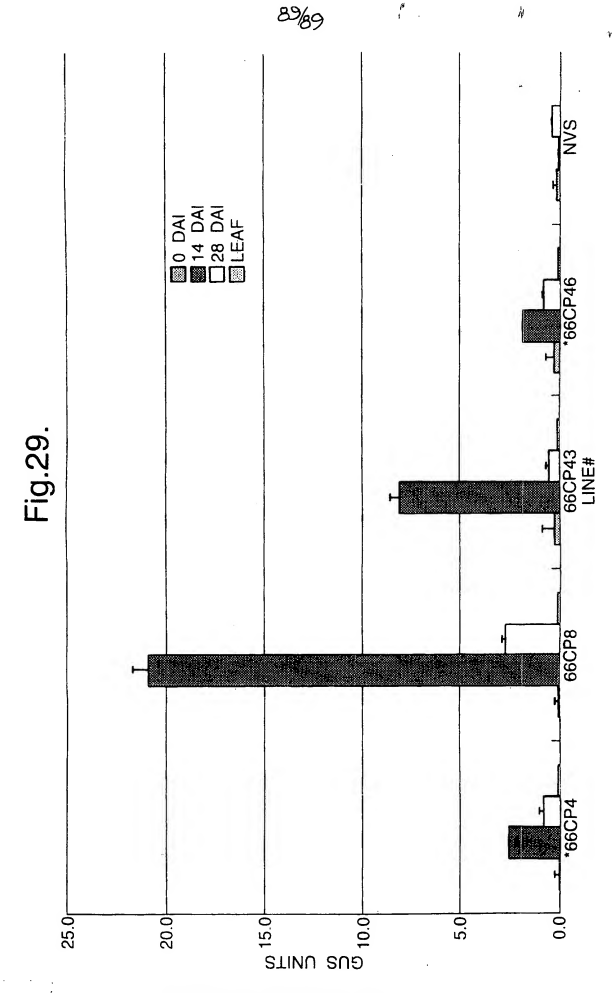


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